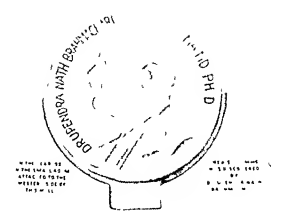
GLEANINGS FROM MY RESEARCHES VOL I KALA AZAR ITS CHEMOTHERAPY



THE CONQUEST OF KALA-AZAR

I PECALL WISH JOY THAT MEMORABLE WENT IN THE CALCULTA CAMPBELL MOSPITAL AT CALCAN WHERE AFTER A VERY HARD DAYS WERK I FOUND AT ABOUT TO DICCOR IN A LITTLE ROOM WITH A SMIRT DIWLY BURN IN LANTERH THAT THE PESULTS OF MY ESPECIMENTS WERE UP TO MY ESPECIATIONS BUT I FIO NOT KNOW THEN THAT PROVIDENCE HAD PUT TITLD MY HARDS A WORDOWS OF THIS DAY THAT THIS LITTLE THING WOULD SAVE THE LIVES OF ANLLINKS OF MY ESPECIATIONS.

I SHALL REVER FORCES THAT ROOM WHERE LUTEA STIRAM HE WAS DISECUTERED THE ROOM WHERE I MAD TO LABOUR FOR MORTH'S WITHOUT A GAS POINT OR A WATER TAP AND WHERE I HAD TO REMAIN CONTENTED WITH AN OLD REPOSENE LAMP FOR MY WORK AT MICHT. THE ROOM STILL REMAINS BUT THE SIGNS OF A LABORATORY IN IT HAVE COMPLETELY DISAPPEARED TO ME IT WILL EVER PERMAIN A PLACE OF PILGRMADE WHERE THE FIRST LIGHT OF UREA STIRAM REDAWNED UPON MY MIND.

TO DAT UREA STIBABURE STANDS PRE EMINERT IN THE TREATMENT OF KALA. AZAR INTROIA AND AS A POWERFUL PROPHYLACTIC AGAINST THE DISEASE AND IT IS A MATTER OF SUPPEME SATISFACTION TO ME THAT THE TREATMENT EVOLVED OUT OF MT RESEARCH HAS REMOVED THE TERRORS OF THIS DISTRESSING DISEASE IT MAY BE HOPED THAT BEFORE CONGITHE DISEASE WILL BE COMPLETELY BAMISHED FROM INDIA AND DTHER PARTS OF THE WORLD WHERE IT DECURS THAT WAL RE THE HAPPIEST AND PROUDEST DAY OF MY LIFE IT IT FALLS TO MY LOT TO SEE IT

(EVTACTE FARM D. BRAHM CH. RES PRET OF EAL. BRACES AT T. CARROLL. MINVERSARY MEETING OF T. C. CLAYE, BOCHTY BY REMEAL. B28.)

GLEANINGS FROM MY RESEARCHES

Vol I KALA-AZAR, ITS CHEMOTHERAPY

BY

SIR UPENDRANATH BRAHMACHARI KT MA MD PHD FRASB FNI

Professor of Tropical Medicine Carmichael Medical College Professor of Biochemistry University of Calcutta

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DEDICATED TO THE MEMORY OF THE POOR MISERABLE SUFFERERS
WHO DIED OF KALA AZAR BEFORE THE DAYS OF ANTIMONY
AND TO THOSE WHO HAVE BEEN SAVED FROM THE
HORRORS OF THIS DREADFUL DISEASE THROUGH
THE LABOURS OF THE AUTHOR IN THE MIDST

OF TROUBLES AND DIFFICULTIES

CONTENTS

	PAGES
Foreword	יוצ וווצ
List of Illustrations	xv xx
A New form of Cutaneous Leishmaniasis-	
Dermal Leishmanoid—By U N Brahma	
charı	1.7
A Note on a new disease- 'Dermal Leishmania	
sis (Brahmachari)—By J W D	
Megaw, Lt Col, IMS	8 10
Chemotherapy of Antimonial Compounds in	
Kala azar Infection-Part/I-By U N	
Brahmachari	11 50
Chemotherapy of Antimonial Compounds in	
Kala azar Infection-Part II-By U N	
Brahmachari	51 55
Chemotherapy of Antimonial Compounds in	
Kala azar Infection-Part III-By U N	
Brahmachan	56 77
Chemotherapy of Antimonial Compounds in	
Kala azar Infection-Part IV-By U N	
Brahmachari	78 92
Chemotherapy of Antimonial Compounds in	
Kala azar Infection—Part V—By U N	
Brahmacharı	93 99
Chemotherapy of Antimonial Compounds in	
Kala azar Infection—Part VI—By U N	
Brahmachari	100 104
Chemotherapy of Antimonial Compounds in	
Kala azar Infection—Part VII—By U N	
Brahmacharı and P Sen	105 108

	Pages
Chemotherapy of Antimonial Compounds in	
Kala-azar Infection—Part VIII—By U.	
N. Brahmachari, in collaboration with	
S C Choudhury, J Das and P Sen	109-119
Chemotherapy of Antimonial Compounds in	
Kala-azar Infection—Part IXBy U. N	
Brahmacharı .	120-135
Chemotherapy of Antimonial Compounds in	
Kala-azar Infection—Part X—By U. N	
Brahmachari, in collaboration with P Sen	136-154
Chemotherapy of Antimonial Compounds in	
Kala-azar Infection—Part XI—By U. N.	
Brahmacharı .	155-165
Chemotherapy of Antimonial Compounds in	
Kala-azar Infection—Part XII—By U N	
Brahmacharı	166-171
Chemotherapy of Antimonial Compounds in	
Kala-azar Infection—Part XIII—By U N.	
Brahmacharı, ın collaboratıon with Major	
Shortt .	172-176
Chemotherapy of Antimonial Compounds in	
Kala-azar Infection—Part XIV—By U	
N Brahmacharı, ın collaboration with	
B B Maity	177-186
Chemotherapy of Antimonial Compounds in	
Kala-azar Infection—Part XV—By U. N	107 100
Brahmachari, in collaboration with J Das	187-190
Chemotherapy of Antimonial Compounds in	
Kala-azar Infection—Part XVI—By U. N	
Brahmacharı, ın collaboration with B B. Maity	191-195
Chemotherapy of Antimonial Compounds in	171-177
Kala-azar Infection—Part XVII—By U.	
N Brahmacharı	196-197

CONTENTS

IX Diana

	I AGES
Chemotherapy of Antimonial Compounds in Kala azar Infection—Part XVIII—By U N Brahmachari in collaboration with	
J Das	198 199
Chemotherapy of Antimonial Compounds in Kala azar Infection—Part I (New Series) —By U N Brahmachari in collaboration	
with I Das and S C Baneriee	200 204
Chemotherapy of Antimonial Compounds in	200 20 1
Kala azar Infection—Part II (New Series) —By U N Brahmachari in collaboration	
with B B Maity	205 206
Chemotherapy of Antimonial Compounds in Kala azar Infection—Part III (New Series)	
—By U N Brahmacharı ın collaboration	
with A M Dutt	207 209
Chemotherapy of Antimonial Compounds in	
Kala azar Infection—Part IV (New Series)	
By U N Brahmacharı and S C Banerjee	210 211
Studies in Kala azar and Chemotherapy of Antimony—Part I—By U N Brahma	
chari and R Banerjea	212 216
Studies in Kala azar and Chemotherapy of Antimony—Part II—By U N Brahma chari in collaboration with J M Das	217 224
Gupta R Banerjea and B N Basu	217 224
Studies in Kala azar and Chemotherapy of Antimony—Part III—By U N Brahma	
chari in collaboration with P B Sur and R Banerjea	225 230
Studies in Kala azar and Chemotherapy of	
Antimony-Part IV-By U N Brahma	
charı	231 232
B767B	

PAGES

	INGES
Studies in Kala-azar and Chemotherapy of Antimony—Part V—By U N Brahma-	
chari, in collaboration with R Banerjea Studies in Kala-azar and Chemotherapy of Antimony—Part VI—By U. N Brahma-	233-236
chari	237-249
Forms of Pyrexia due to Leishman-Donovan Bodies—By U N Brahmachari	250-251
A Contribution to the Study of Fevers due to Leishman-Donovan Bodies—By U N	
Brahmachari	252-260
Transactions of the Calcutta Medical Club	261-264
Sporadic Kala-azar in Calcutta, with Notes of a case treated with Atoxyl—By U N	
Brahmacharı	265-273
Fatty Liver in Kala-azar—By U N Brahma- chari	274-275
A Preliminary Report on the Treatment of Kala-azar with Intravenous Injection of Metallic Antimony—By U N Brahma- chari	276-281
Further Observations on the Treatment of Kala- azar and cases treated with Metallic Antimony, Sodium Antimonyl Tartrate, Formaldehyde and other Drugs—By U N Brahmachari	282-290
Third Report on the Treatment of Kala-azar with Special Reference to the Use of Antimony and Formaldehyde—By U N.	201.004
Brahmacharı Envil Brahmacharı	291-304
Fourth Report on the Treatment of Kala-azar and some Blood Reactions in this Disease —By U N Brahmachari	305-314

	Pages
On the Presence of an Easily Precipitable Anti-complementary Globulin like Sub- stance in Human Serum and its Import ance in the Diagnosis of Kala azar—By U N Brahmachari	315 321
Treatment of Kala azar with Intramuscular Injections of Hyper acid Antimonyl Tartrate (and Urethane)—By U N Brahmachari	322 325
A Prehminary Note on the Globulin Albu min and Cholesterol Contents of the Blood in Kala azar—By U N Brahma chari and M M Dutt	326 328
The Freatment of Kah azar with some New Antimonial Preparations—By U N Brahmachani	329 3 37
The Globulin Opacity Test in Kala azar— By U N Brahmachari and P B Sen Urea Stibamine in Kala azar—By U N Brahma	338 342
chari The Relation between the Chemical Constitution of Antimonials and their Thera	343 352
peutic Properties—By U. N. Brahmachari Proceedings of a Meeting held at the Calcutta	353 373
Medical Club on the 14th July 1916 Preparation of Urea Antimonyl Tartrate a	374 380
New Compound—By U N Brahmacharı A Contribution to the Chemistry of certain New Aromatic Antimonials—By U N	381
Brahmacharı and J M Das Gupta Synthesis of a few Antimonials of Therapeutic Interest—By U N Brahmacharı and	382 388
J M Das Gupta	389 394

	Pages
Synthesis of Sodium <i>n</i> -Phenyl-glycine-amide-	
4-stibinate (Antimony Analogue of	
Tryparsamide)—By U N Brahmachari	395
The Intensive Antimonial Treatment of Kala-	
azar—Part I—By U N Brahmacharı	
and P N Brahmacharı .	396-401
The Intensive Antimonial Treatment of Kala-	
azar—Part II—By U N Brahmacharı,	
A R Mazumdar and R B Dey	402-419
Campaign against Kala-azar in India—By U N	
Brahmacharı	420-427
Chemotherapy of Antimonial Compounds—By	
U N Brahmacharı	428-433
Urea Stibamine in the treatment of Indian Kala-	
azar—By U. N. Brahmacharı	434-440
The Conquest of Kala-azar—By U N Brahma-	
charı	441-452

FOREWORD

The author's apology for reprinting some of his published papers in a collected form is to infuse the spirit of research into the minds of students of medicine in India and not least amongst those whose paths are restricted to institutions where proper facilities for research are not yet available. These collections also create a sense of happiness in the mind of the author as they remind him of his own past struggles in carrying on his researches. There has been no attempt to observe a logical or even a chronological sequence and the papers are set out in the manner which it is hoped is best calculated to provide stimulating and attractive reading

The first volume of this work contains a series of the author's papers on kala azar including chemotherapy of antimonial compounds in kala azar infection, which have appeared from time to time in various journals They record the evolution of his advances in the treatment of a terrible tropical disease. The more ambitious student will find some portion of the subject dealt with at greater length in the author's previous works on kala azar such as (1) Kala azar Its Treatment (Butterworth & Co Calcutta 1917) and subsequent editions (2) Treatise on Kala azar in German Professor Dr Carl Mense's Handbuch der Tropen krankheiten Vol II 1926 and (3) A Treatise on Kala azar (John Bale Sons & Danielsson Ltd London 1928) these will be found many conclusions which the author has formed after continued research, but which have not been the subject matter of previous articles and are therefore omitted from the present work. The last chapter in this volume The Conquest of Kala azar is a portion

of the author's Presidential Address at the Medical Section of the Indian Science Congress, Twenty-fifth Session, 1938.

The second volume contains the author's papers on malaria including his studies on the chemistry and chemotherapy of quinoline and actidine compounds which resulted in the synthesis of a compound allied to atebrin The first chapter in this volume is a continuation of the author's Presidential Address, referred to above It consists of certain observations on the chemotherapy of malaria.

It is intended that the author's other published papers will form the subject matter of another volume

Some of the articles in these volumes have been edited by the writer

The author hopes that a time may come when his past struggles in research will see the light of day

LIST OF ILLUSTRATIONS

Plates

Faci	ng Pa
The different organs cut open showing extensive hæmorrhages into their substance brought about by	
antimony poisoning Plate XII	14
Section of lung showing hamorrhage into the intersti	
tial tissue Plate XIII	14
Section of lung tissue showing hæmorrlinge round cell infiltration and blocking of alveoli with debris	
Plate XIV	14
Section of Lidney showing hæmorrhage into the inter stitial tissue cloudy swelling and destruction of the Lidney epithelium and exudation of granular	
material into the kidney tubules Plate XV	14
Section of liver showing round infiltration round the portal system fatty degeneration cloudy swelling	
and hæmorrhage Plate XVI	15
Section of spleen showing destruction of splenic pulp Plate XVII	15
Section of suprarenal gland showing hæmorrhage in	
zona reticularis and medulla Plate XVIII	16
Section of pituitary pars anterior normal—Fig 1 Plate XIX and showing diminution of eosinophile	
staining contracted appearance of cells and	
hæmorrhage-Fig 2 Plate XIX	16
Section of pituitary pars posterior normal-Fig 1	
Plate XX, and showing marked diminution of	
eosmophile staining of the cells, nuclei of the cells	t
contracted Death from tartar emetic poiloning	
the animal died seven days after injection—Fig 2	
Plate XX	16

Facing	g Page
Section of pituitary pars anterior showing marked diminution of eosinophile staining of the cells, the nuclei of the cells contracted, and interstitial hæmorrhage. Death took place seven days after	
emetic injection Plate XXI	16
Section of parotid gland showing degeneration of glandular cells, and hæmorrhage. Plate XXII	16
Photograph of a patient—first case of Dermal Leishmanoid (vide p 4 also), showing the cruptions on the upper part of the patient's body Plate	
LXXIII	51
Scrapings from one of the papules, showing the presence of L D bodies which are mostly free forms and extracapsular, a few are inside leucocyte and endothelial cells Fig. 1, Plate LXXIV Vide	
also p 53	52
Scrapings, showing flagellated bodies found after 12 days which were indistinguishable from those of	
L D bodies Fig 2, Plate LXXIV Photograph of a monkey, showing marked granulo- matous growths at the site of inoculation in both eyebrows, done a month and a half before, by embedding bits of granulomatous nodules into pockets cut in them, with secondary nodules at the	52
outer and inner canthuses of the eyes Plate LXXIV (b) Fig. 3, Plate LXXIV shows a fair number of L D. bodies in smears from one of the incised nodules taken from the original site of inoculation made in the monkey represented by Plate	52
LXXIV (b) Fig 4, Plate LXXIV, section of a papule in the skin of the patient showing round-celled infiltration	52
with fibroblasts and thinning of the epidermis	52

52

	no Pa
Fig 5 Plate LXXIV—the same as Fig 4 showing the presence of a network of newly formed	
capillaries and thickening of capillary wall. Photograph of a patient showing peculiar eruption of	52
the body nine months after completion of the second course of treatment—a case of Dermal Leishmanoid Vide para 2 Plate XXXII (Vide	1770
also p 208)	173
Section showing appearances of epidermis and culi vera in superficial tissue taken from the nodules of a Dermal Leishmanoid case under a low power	
Fig 1 Plate XXXIII para 2	175
Section showing appearance of epidermis and cutis vera in superficial tissue from nodule of a Dermal Leishmanoid case under high power Fig 2 Plate	
XXXIII para 5	175
Photograph of a case of Dermal Leishmanoid supposed to have been a refractory case but subsequently cured by persistent treatment with urea stiba	
mine Plates 1 and 2	213
Photomicrograph of spleen of a mouse infected with Leishmania Donovani forty eight hours after intravenous injection of metallic antimony—	
A—Cell containing leishmania but no particles of antimony	
B—Cell with faintly stained cytoplasm containing leishmania and a few particles of antimony	
C—Cell containing coarse granules of antimony and leishmania some of which appear to be degenerated Para 6	226
Photomicrograph of spleen of a mouse infected with	
Leishmania Donovani forty eight hours after	
intravenous injection of metallic antimony	232

C-767B

Fac	ing Page
Section of liver showing extensive fatty degeneration of the cells, more about the peripheral portions of the lobules of the liver Para 3	
$D_{lagiams}$	
Diagram of a curve formed showing the values obtained for the M L D and M T D of various antimonyl tartrates <i>Vide</i> p 27, also Plate XXIII Chart showing the curve of excretion of antimonyl	23
tartrates after intravenous injection in terms of antimony (Chart 1)	115
1 and 2 Curves of excretion of tartar emetic in terms of antimony after repeated injections of tartar emetic (2 cases)	
A theoretical curve showing the excretion of tartar emetic in terms of antimony after repeated injec-	
tions if the rate of excretion always followed the law that the amount excreted was proportional to the amount present in the system .	115
Excretion of antimony after injection of tartar emetic and urea stibamine	136
Diagram of a special syringe and equipment used for author's own method of injecting metallic antimony intravenously	277
Temperature Charts	
Temperature chart of a kala-azar case treated successfully with urea stibamine	75
Temperature chart of a kala-azar case showing high intermittent pyrexia (Chart 1)	250
Temperature chart of a case showing irregular intermittent pyrexia (Chart 2)	250

Face	ng Page
Temperature chart of a kala kazar case showing	
Double quotidian pyrexia with double intermissions	
during 24 hours (Chart 3)	250
Temperature chart of a kala azar case showing	
Double quotidian pyrexia with single intermission	
in 24 hours (Chart 4)	250
Temperature chart of a kala azar case showing	
Double remittent pyrexia (Chart 5)	251
Temperature chart of a kala azar case showing	
combined intermittent and remittent pyrexia resem	
bling hectic (Chart 6)	251
Temperature chart of a kala azar case showing an	
almost apyretic temperature (Chart 7)	251
Temperature chart of a kala azar case treated with	
atoxyl Para 3	270
Chart showing effect of hetol on leucocyte count	270
Temperature chart of a kala azar case treated with	270
electrargol and metallic antimony Case No 4 Temperature chart of a kala azar case treated with	278
tartar emetic antimonyl sodium tartrate and	
metallic antimony Para 2	278
Temperature chart of a kala azar case first treated	270
with tartar emetic and plimmer's salt then after	
some days treated with intravenous metallic	
antimony Para 2 (ref p 286)	278
Temperature chart of a kala azar case treated with	
electrargol and tartar emetic Case No 6 Table	
on p 281	278
Temperature chart of a kala azar case treated with	
three injections of metallic antimony half a grain	
each time Para 1, Case No 2	283
Temperature chart of a kala azar case treated with	
two injections of metallic antimony Case No	20.4
5 K	284

Amiyo. Chart (a)

Facin	ng Pag
Temperature chart of a kala-azar case treated with	
antimony sodium tartrate after berberine failed,	
vide Chart (a), p 303, para 2, subsequently	
cured by colloidal metallic antimony, patient-	
Amıyo Chart (b) .	292
Temperature chart of a kala-azar case treated	
first with intravenous narcotine with no improve-	
ment in temperature, subsequently treated with	
metallic antimony Case No 2, patient—Abdul	
Azız, ref p 302	294
Temperature chart of a kala-azar case treated with	
three injections of formaldehyde with remarkable	
ımprovement Case No 2, patient—Tetarı	300
Temperature chart of a kala-azar case treated with	
berberine sulph intravenously but discontinued as	
there was no improvement. Para 2 nations—	

303

A NEW FORM OF CUTANEOUS LEISHMANIASIS-DERMAL LEISHMANOID

The following paper on A New Form of Cutaneous Leishmaniasis was read by me at the meeting of the Medical Section of the Asiatic Society of Bengal held on 8th February 1922

The various forms of cutaneous and muco cutaneous leishmaniasis are divided by Castellani and Chalmers as follows —

- (1) Cutaneous
- (2) Muco cutaneous
- (3) Oro pharyngeal

The cutaneous forms are divided by them into -

- (a) The common variety—The oriental sore
- (b) The verrucose variety
- (c) The keloid form
- (d) The frambœsiform
- (e) The papillomatous variety
- (f) The deep ulcerative variety

Laveran describes the following forms of cutaneous

- (A) The oriental sore
- (B) American leishmaniasis (a) the cutaneous ulcerating form (b) the cutaneous non ulcerating form which may be either papillomatous or macro tuberculous

The variety of cutaneous leishimaniasis described in the present paper is of extreme pathological and clinical importance. It differs from any form of cutaneous leishimaniasis described in literature and appears to afford the missing link between cutaneous and visceral leishimaniasis or kala-azar and leads one to conclude that the special pathogenic properties of the parasites of kala-azar may be so modified after antimonial treatment that it may subsequently give rise clinically to a form of cutaneous leishimaniasis, thus proving the identity of the parasite of kala-azar and that of cutaneous leishimaniasis.

Among the multitude of kala-azar patients treated by me with intravenous injection of antimony, I met with four cases which, within six months to two years after completion of treatment, came to me with a peculiar form of cutaneous eruptions which, at first sight, gave an impression of tuberculous leprosy. In none of them, however, could any lepra bacilli be found. When they came to me with these eruptions, there were no clinical symptoms of kala-azar.

The appearance of these cutaneous eruptions in patients who have apparently recovered from kala azar after antimonial treatment made me suspect that they might be due to a cutaneous infection of these individuals in whom there was not a complete sterilization of the organs against the leishmania, though their virus had been attenuated by repeated antimonial injections. This led me to examine the scrapings from the cutaneous nodules of these cases with the help of Dr. Surendra Nath Ghose, Bacteriologist, Presidency General Hospital, Calcutta. The examination of the scrapings led to the remarkable discovery that the eruptions were due to cutaneous infection by the parasites of kala-azar.

During the antimonial treatment of kala-azar, the following results may follow —

- (1) Cure
- (2) Apparent cure followed by a relapse
- No improvement

A fourth result may follow and this is what happened in the four cases mentioned above. The visceral leish maniasis may be cured but a few leishmania may be left behind with their virus so attenuated that they gave rise to a milder disease namely cutaneous leishmaniasis.

l give here the full history of the last case in which this transformation of a case of visceral leishmaniasis (kala azar) into one of cutaneous leishmaniasis took place

Patient æt 31 nn inhibitant of Barisal gave a history of fever coming on with rigors from February 1917 which was not benefited by quinne In May 1917 he had an attack of pneumonia. His fever persisted and there was progressive enlargement of the spleen. He was again treated with quinne which was given intramuscularly in doses of 10 grains for 6 days. He states that after this he was free from fever till the end of June 1917. In July he again had an attack of intermittent fever, the temperature ranging between 99° F and 105° F. He was again given intramuscular injections of quinne but with no benefit.

In January 1918 he came to Calcutta for the treatment of persistent fever with the enlargement of the spleen and the liver. The spleen extended 6 inches below the costal margin and the liver extended 3 inches below the costal arch. The fever was of an intermittent type. He was at first given a course of treatment with soamin. The results of blood examination before treatment with soamin were R. B. C. 3.000.000 W. B. C. 3.500. Hb. 30 per cent and differential count showed polymor phonuclears 60 per cent. Lymphocytes 24 per cent. large mononuclears 14.8 per cent. and eosinophiles 1.2 per cent.

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The treatment with soamin was not followed by any improvement Spleen puncture was made and the smear showed the presence of Leishman-Donovan bodies. A few L D bodies were also found in peripheral blood. The patient was now treated with intravenous injection of tartar emetic given twice a week in doses of ½ to 10 c c. He had altogether thirty injections. The fever stopped after 10 injections. When he left the treatment, there was marked improvement in his general condition, the spleen and the liver could not be felt below the costal margin and the blood condition was—

R B C 4,000,000 W B C 7,500 Hb 70 per cent

No parasites could be found on spleen puncture

He has had no fever since his treatment with antimony was stopped

In the beginning of 1919, he noticed faint whitish patches on his face. These gradually spread. These patches were neither anæsthetic nor hyperæsthetic. They gradually spread over the whole body in front and behind in about six months. He was at first treated with arsenic internally. The patches became worse during cold weather. Subsequently, papillomatous nodules appeared over the face, the trunk and the extremities

Patient was seen by me very recently I asked Dr Ghose to make a very careful examination of the scrapings and the juice from the papillomatous nodules for the presence of L D bodies. The smears showed a very large number of L D bodies in some of the slides.

Description of the present rash — The whole of the body is covered with eruptions which are described as follows —

- (1) On the face there are papillomatous nodules somewhat resembling small leprotic nodules
- (2) There is a slight erythematous appearance on the cheeks and the forehead

- (3) On the trunk, the upper and the lower extremities there are slightly raised brown patches which are extensively spread over the whole body A few papules are also present in these parts
- (4) There are some erythematous patches in the extremities especially the lower
- (5) No ulceration or scab formation in any part of the body. Other features—no anæsthesia no loss of knee jerks no thickening of the nerves. No eruptions in the mucous membrane of the mouth and nostrils.

Liver and spleen normal On examination of the splenic blood by spleen puncture no L D bodies were found No rise of temperature. The patient complains of no other trouble except the ugly appearance of the body due to the eruptions.

Result of blood examination on 1st February 1922 -

Hb 75 per cent
R B C 4 500 000
W B C 10 000
Polymorphonuclears 62 per cent
Lymphocytes 24 per cent
Large mononuclears 6 per cent
Eosnophiles 8 per cent

The blood report does not at all correspond to that of Lala azar No L D bodies could be detected in the peripheral blood

Examination of the scrapings—L D bodies are found in very large numbers especially in the juice expressed from the papillomatous nodules A few have also been found from the brownish patches No lepra bacilli

In view of the fact that the eruptions are due to leishmania infection whose virus has been modified by antimonial treatment. I propose to call this form of cutaneous leishmaniasis dermal leishmanoid just as small pox modified by vaccination is called varioloid.

I shall study the morphological character of the flagellate forms of these parasites after culturing them with the help of Major Knowles, I.M.S., Protozoologist, Calcutta School of Tropical Medicine.

This case, along with three others of a similar type that I have observed, is a remarkable one, as they appear to point to the identity of the parasites of visceral and cutaneous leishmaniasis

It seems that the virus of the parasite of kala-azar was attenuated in these cases by the antimonial treatment and a case of deadly visceral leishmaniasis was converted into one of cutaneous leishmaniasis. We thus have a direct proof of the identity of the parasites of visceral and dermal leishmaniasis, which has been attempted to be proved indirectly by complicated inoculation experiments

Of the three other cases met with by me, one resembled the present case, the rash being generalized over the whole body. The other two cases had less generalized rash, most of the papillomatous eruptions being present on the face, there being some brownish patches over the arms

One of these cases was treated with further injections of antimony and he appeared to improve The second one, a boy of 15 years, was given six intravenous injections of tartar emetic in doses of 3 to 5 c c, but he left treatment before any improvement was noticed I propose to treat the present case with combined treatment of intravenous injection of antimony and soamin and shall report the results in a future communication

It has been suggested by Manson that the treatment of kala-azar with a vaccine made from the virus of oriental sore is worth trial. May it be further suggested that in places where kala-azar is very prevalent, the inhabitants should be vaccinated with the virus of oriental sore as a prophylaxis against kala-azar?

Apart from the interest in the above case on account of its forming a new hitherto unknown clinical entity it raises the following most suggestive questions —

- (1) Are the parasites of kala azar in the process of destruction by antimonial treatment eliminated by the skin and are cases of kala azar therefore more infective during antimonial treatment?
- (2) If the parasites are eliminated by the skin do they also enter the system through the skin at the time of primary infection?

The above case after being exhibited by me at the meeting of the Medical Section of the Asiatic Society of Bengal held on 8th February 1922 was exhibited at the Calcutta School of Tropical Medicine on 9th February 1922

l append here a drawing showing the eruptions on the upper part of the patient s body. A drawing from the scrapings from one of the nodules is also appended herewith showing the presence of Leishmania donovani which mostly seems to be extra corpuscular in the smear. As stated before I have met with four cases of dermal leishmanoid.

Perhaps such cases are more common than has been suspected and more cases will be met with by observers who are treating kala azar with antimonal preparations

l am indebted to the Editor Indian Medical Gazette for announcing my discovery of this new form of cutaneous leishmaniasis in the Indian Medical Gazette for March 1922

I suggest that workers in the field of kala azar should look out for such cases of infection by Leishmania donovani sine kala azar as a result of antimonal treatment

Since the above paper was sent to the editor Indian Medical Gazette 1 have succeeded in developing flagellated forms of Leishmania donovani with the help of Major R Knowles IMS on NNN medium from the juice obtained from the eruptions by puncture Blood cultures were negative

A NOTE ON A NEW DISEASE—"DERMAL LEISHMANIASIS" (BRAHMACHARI)

By J W D MEGAW, LIEUTENANT-COLONEL, I M S

The above paper on "dermal leishmanoid" by Dr Brahmachari is of very exceptional interest. It records a disease hitherto unknown to science, but what is more remarkable is that the disease appears to have been produced by human agency. As the writer of this note remarked when Dr Brahmachari showed one of the cases at the Calcutta School of Tropical Medicine, it is a unique experience for a medical man to be the agency through which a new disease is produced

The disease differs entirely from drug eruptions in being a specific parasitic affection which follows the administration of antimony salts by the intravenous route

It is possible that the disease may occur under other conditions but there is no evidence that this takes place, and it is not likely that a dermal leishmaniasis of so distinct a kind would have escaped notice hitherto, considering the number of practitioners who are in the habit of examining scrapings in all doubtful cases of chronic skin disease. Those who have seen the case shown by Dr Brahmachari, agree that the disease is a hitherto unknown form of leishmaniasis and Dr Brahmachari is to be heartily congratulated on the very interesting and important discovery which he has made

There will be differences of opinion as to the interpretation of the findings. It has not yet been proved that the cutaneous leishmaniasis is due to a modified virus

C

There is some factor which has caused the parasite to behave in so remarkable a manner but it remains to be seen whether the virus is changed in its nature. Indeed the very example which is given by Dr. Brahmachari. viz varioloid is really an example of virus whose nature is not essentially changed. It is rather the human body which reacts differently in cases of varioloid and the same thing probably happens in the case of the dermal leishmaniasis under discussion.

There is however an important difference in that the parasite in dermal leishmaniasis produces manifestations which are essentially different in type from those of kala azar of which it is an after development—It is also impossible to agree with Dr. Brahmachari when he says that the case appear to point to the identity of the parasites of visceral and cutineous leishmaniasis. What the cases point to is the identity of the parasites of this particular form of dermal leishmaniasis with those of kala azar but it does not follow that the parasites of other forms of leishmaniasis such as criental sore are identical.

The parasites from one of Dr Brahmachari's cases have been cultivated by Major Knowles and as would be expected the cultures appear to be identical with those of the Leishman Donovan body and up till the present there is no evidence that they are in any way different from that parasite. Meantime it is to be hoped that other workers will look out for further cases of this disease especially in Assam where thousands of cases of kala azar have undergone treatment by antimony preparations

A look out should of course be kept for possible cases arising independently of antimony treatment

Already another case of the disease has been recognised by my assistant Dr Bhattacherji in a patient who had under gone antimony treatment

The name proposed by Dr Brahmacharı does not seem to be suitable the disease is a leishmaniasis not a leishmanoid

The name which appears to me to be most appropriate has the serious drawback of being clumsy, it is "Post-Antimonial Dermal Leishmaniasis" (Brahmachari) or more briefly "Brahmachari's Dermal Leishmaniasis" The afterhistory of the cases will be watched with the greatest interest, and meantime there will doubtless be a good deal of speculation as to the reason for the production of the disease

Have the parasites in the accessible parts of the circulatory system been killed off? And have those which remain in the less accessible parts of the circulatory system been cut off from the destructive action of the antimony and the blood by an obliteration of the capillaries as a result of the action of the antimony? If such an action results from the antimony injections, it is possible that it is a question of the parasites being entrapped in places where they are capable of growing unhindered by whatever immunising mechanism there is, and that a general immunity against the parasites has no chance of developing, because the parasites in the body in general have been killed off by the antimony

It now remains to be seen whether a course of antimony treatment will influence the skin manifestations and also whether the disease will remain localised in the skin

If any of the cases refuse to undergo antimony treatment there will also be an opportunity of seeing the natural evolution of the disease

[NB —Subsequent history of this case is described in the Transactions of the Royal Society of Tropical Medicine, Vol XXIII, No 3, pp 301-04, issued November, 1929 -Editor 7

CHEMOTHERAPY OF ANTIMONIAL COMPOUNDS IN KALA AZAR INFECTION

[Redfr Puble to Jn 193 1972]

PART I

THE TO\ICITY OF ANTINIONYL TARTRATES—THE INFLU
ENCE OF THE BASIC RADICLE OF AN ANTIMONYL
TARTRATE UPON ITS TO\ICITY—SONIE ARYL PEN
TAVALENT ANTIVIONIAL CO\IPOUNDS—P AMINO
PHENYL STIBINIC ACID AND SO\IC OF ITS
DERIVATIVES—THEIR TO\ICITY—THE
THERAPEUTIC VALUE OF AMMONIUM
ANTIMONYL TARTRATE AND
UREA STIBAMINE

The Symptoms of Antimony Poisoning in the Guinca pig and the Rat after Injection of Antimonyl Tartrates

The symptoms of acute antimony poisoning in the guinea pig and the white rat are not always constant. Vomiting and purging which are frequently observed in man after intravenous injections are not at all common in guinea pigs even after toxic doses and it is not rare to find solid fæces in the large intestines even when the animal dies of severe acute symptoms

The symptom complex of the intoxication produced by the various antimonyl tartrates enumerated below does not vary much whatever may be the salt used. They are divisible into two groups of phenomena.—

(1) Nervous (2)

(2) Nutritional

In some severe cases of acute poisoning, the animal passes into a state of prostration with complete paralysis of the central nervous system within $\frac{1}{2}$ to 1 hour after injection severe cases the symptoms come on more gradually Marked tremor and chattering of the teeth are sometimes very characteristic features of intoxication and then the animal passes into a state of coma and in the comatose state may develop spasmodic movements of groups of muscles coming on at intervals of ½ to 1 minute In fatal cases the breathing is hurried and pulse quick In some cases, the animal lies in a comatose condition for some hours before death. Sometimes the animal shows marked muscular tremors and incoordination when disturbed Salivation has been observed in many cases but is not a constant symptom In some cases a few minutes after intravenous injection the animal exhibits a very marked spasmodic movement of the whole body at frequent intervals In some cases spasmodic contraction of diaphragm resembling hiccough in man has been observed. In many cases, soon after the injection of a fairly large dose, the animal frequently scratches its mouth with its front legs Sometimes even after sublethal doses the animal appears to be ill and faint for a short time It is unsteady, the gait is staggering and the animal may roll about The animal is less active and takes its food less freely than usual. If it survives for 10 or 12 hours after the injection, then there is development of a peculiar bloated appearance of the face in fatal cases In cases that survive, this phenomenon is slightly or not at all marked

There may be marked emaciation in cases that survive 2 or 3 days after injection but frequently the animal regains in weight in a week's time

In some cases, the animal progressively loses in weight It takes very little food, remains dull and dies on the 7th or 8th day. Such cases are rare and generally it may be stated that if recovery takes place, it comes on within two or three days and is complete.

Period at which Death Takes Place after Doses within the Toxic Range

The earliest period at which death took place in guiner pigs after minimum lethal doses was 4 hours. In some cases the animals took 12 to 18 hours to die after injection of the minimum lethal doses. With smaller doses but still within the toxic range the animals that did not survive died 24 to 36 hours after injection. Sometimes the animals survived for 8 to 10 days, and after death they showed symptoms of anitmony poisoning. Cases of delayed antimony poisoning will be described in another series.

PATHOLOGY OF THE INTOXICATION

The pathological changes produced in the animal may be studied under the following heads —

- (1) Local effects
- (2) Systemic effects

It may be stated that generally speaking the local effects produced after intramuscular injections of the antimonyl tartrates are much less marked in the case of the guinea pig and the rat than in the case of man. There may be some irritation and swelling at the seat of injection but necrosis or destruction of tissues is rarely met with—a phenomenon frequently met with in the case of map.

Systemic Effects

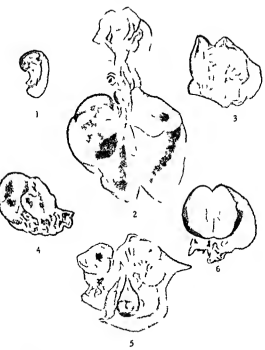
The effects produced by the antimonyl tartrates upon the animal as a whole when given in toxic doses are very marked. In general after toxic doses the pathological lesions consist of hemorrhages into the internal organs and necrosis of their cellular elements. The organs in which the changes are most marked are (1) Lungs (2) Kidneys and (3) Liver. Marked changes in the gastro intestinal tract are

not frequently met with Sometimes there is congestion of or even hæmorrhage or ulceration in the gastric mucosa. There may be hæmorrhages into the substance of the spleen Among the ductless glands that have been studied marked pathological changes may take place in the adrenals and pituitary. No change has been observed in the thyroid

l shall now describe these changes in the different organs in detail —

- of acute poisoning, the whole lungs are in a state of extreme congestion with hæmorrhages into their substance and alveoli. On section, blood pours out freely from the cut surfaces. On microscopic examination, extensive hæmorrhages into the substance of the lungs with destruction of the parenchyma with round-celled infiltration and exudation of necrosed material into the alveoli of the lungs are met with (See Plates XII, XIII and XIV) In one animal there was evidence of lobar pneumonia in one lung, but this might have been accidental
- (2) Kidneys —In fatal cases, marked destructive changes are met with in the kidneys. In acute cases, the kidneys are slightly enlarged. There may be hæmorrhages into their capsules. The congestion is sometimes most marked in the boundary zone and frequently extends outwards along the medullary rays towards the capsular surface. Sometimes there is cloudy swelling and sometimes necrosis of the kidney epithelium. Hæmorrhage which may sometimes be very extensive may be seen in the interstitial tissues of the kidney. The tubules of the kidneys may be blocked with granular debris. (See Plates XII and XV.)
- (3) Liver—In some cases, the liver presents a pale, yellowish appearance which is indicative of extensive fatty change. On the surface there may be spots of hæmorrhage, sometimes very extensive. In other cases the liver presents a deeply congested appearance with hæmorrhages into its

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1-SPLEEN

2-LUNGS

3-STOMACH [opened]

4-KIDNEY

5-GALL BLADDER 6-KIDNEY [pen d]

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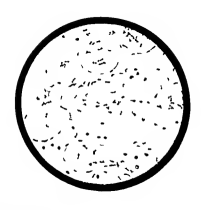
PLATE XIII



Sect on of lung how ng hæmorth g into the te tit al tissue

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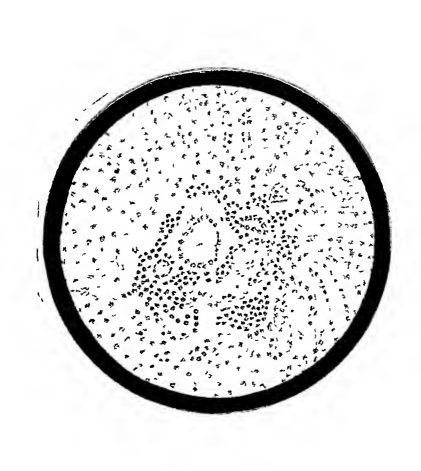
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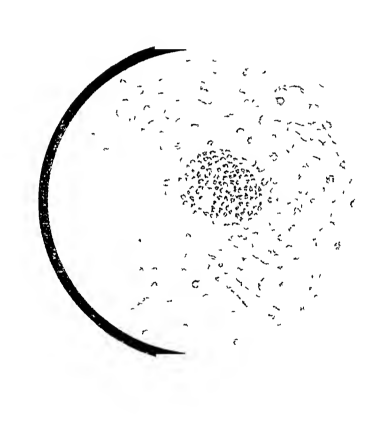
PLATE XVI



Section of Liver showing round infiltration round the portal system, fatty degeneration, cloudy swelling and hæmorrhage

[Reprinted from the Indian Journal of Medical Research Vol X, No 2, October, 1922]

PLATE XVII



Section of Spleen showing destruction of spleen pulp

substance The latter is observed in those cases in which the animal dies within a few hours after injection and the former when the animal dies after more than 24 hours after injection. There may be hæmorrhages into the substance of the gall bladder. Bile may be blood stained. (See Plate XII.)

On microscopic examination the following changes are noticeable —(See Plate XVI)

- (a) Round celled infiltration around the portal system and hæmorrhage into the interstitual tissue
- (b) Necrosis and extensive fatty degeneration of the hepatic cells
- (c) Blocking of bile capillaries with granular debris
- (4) Spleen—There may be hæmorrhages into the sub stance of the spleen Necrosis of the splenic pulp may be observed (See Plate XVII)
- (5) Gastro intestinal Tract—In some cases there may be signs of acute congestion with patches of ulceration in the stomach but this is not constantly met with Sometimes the small intestines are deeply congested and there may be hemorrhages into their peritoneal coating. The large intestines frequently escape and there may be solid fæces niside them.
- (6) Salwary Glands —There may be extensive destruction of the secreting cells of the parotids with hæmorrhage and round celled infiltration (Plate XXII)
- (7) Ductless Glands —(a) Thyroid No changes have been observed in the thyroid
- (b) Adrenals Hæmorrhages into the substance of the adrenals are not infrequently observed in the acute cases. The cortical vessels may be swollen There may be marked

decrease in the cortical pigmentation Degenerative changes may be observed in the cortex and medulla. (Plate XVIII)

- (c) Pituitary The changes in the pituitary may be divided into two classes --(See Plates XIX, XX and XXI).
 - (1) Changes that take place in the gland after death from severe acute poisoning
 - (11) Changes that take place in the gland after death from subacute poisoning

In the former, there may be hæmorrhages into the substance of the anterior portion of the pituitary with diminution in the eosinophile staining of the cells. In the latter, marked increase in the basophile staining of the cells with slight hæmorrhages is the characteristic change and there may be shrinking of the protoplasm and the nuclei of the cells with prominence of the interstitial tissue. The same changes may be more or less present in the posterior portion of the pituitary.

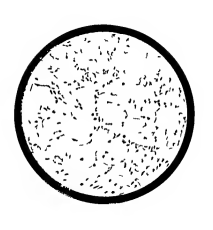
The Toxicity of Antimonyl Tartrates—The Influence of the Basic Radicle of an Antimonyl Tartrate upon its Toxicity

Since the discovery of antimony as a specific in the treatment of leishmaniasis, no systematic work has been done to determine the toxicity of the antimonyl tartrates. It is at the same time evident that such an investigation should be of the highest importance as the antimonyl tartrates are the compounds that are still most commonly used in the treatment of the various forms of leishmaniasis

In the present paper, the toxicity of the following antimonyl tartrates has been investigated —

(1) Ammonium antimonyl tartrate, (2) Urea antimonyl tartrate, (3) Aniline antimonyl tartrate, (4) Potassium antimonyl tartrate, (5) Sodium antimonyl tartrate

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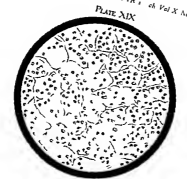
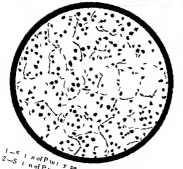
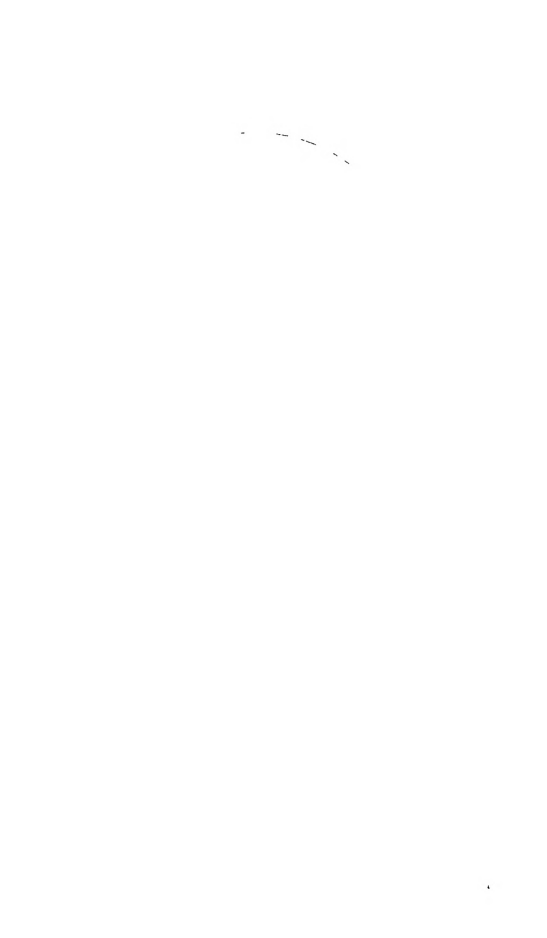


FIG 1



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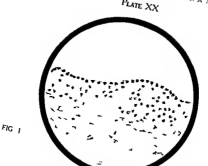
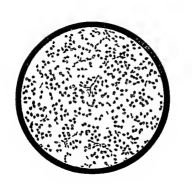




FIG 2

Fic 1-Sect n fPtu mprp t Fig 2—Set not Ptutty p Potto Dath f m it mip nog the an malddyn dy ft inj ton d_{m nut} n f tnhwmrkd nucl of th || noph I t n ng of th

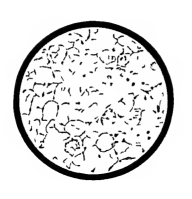




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UREA ANTIMONYI, TARTRATE

In the process of my chemotherapeutic investigations I have sucreeded in preparing a new antimonyl tartrate the urea antimonyl tartrate the preparation and properties of which have been de cribed by me in the Journal and Proceedings Asiatic Society of Bengal (New Series Vol XVI 1920 No 8) The therapeutic value of this antimonyl compound in kala azar has been recorded by me in the Journal of Tropical Medicine and Hygiene August 15 1921 Since then with the help of one of my assistants I succeeded in preparing this compound in another way which is described as follows—

I gram of urea is gently heated with an aqueous solution of 5 grams of tartaric acid for about half an hour. This gives a solution of acid urea tartrate which is subsequently concentrated by gentle heating. To the concentrated solution of the acid urea tartrate a small weighed quantity of Sb O_i is added and the mixture gently boiled till the Sb O_i goes into solution. This process is repeated till 4.8 grams of Sb O_0 are dissolved. The solution is then filtered and concentrated to a syrupy consistency and then allowed to crystallize. In 24 to 48 hours, beautiful crystals separate which are removed and dried on a porous plate and purified by repeated crystallization. Yield ≈ 8 grams

The salt originally prepared by me corresponded to the following formula — CO(NH) C₄H₅SbO O₆ 5H₂O Prepared in the above way it contains ¹HO as water of crystallization

COMPOSITION

Calculated for CO(NH) (C₃H SbO O₈) ¹H O Sb=37 55 ¹N N=4 38 ¹N C=16 9 ¹N H=2 34 ¹N Found Sb=37 55 ¹N N=4 28 ¹N C=16 8 ¹N H=2 2 ¹N H=2 It thus appears that in urea antimonyl tartrate, urea combines with two equivalents of antimonyl tartaric acid, being therefore different from other salts of urea, in which only one of the amino groups in urea is neutralized by the carbonyl group

AMMONIUM ANTIMONYL TARTRATE

It is best prepared by the interaction of acid ammonium tartrate with Sb_2O_3

67 grams of acid ammonium tartrate mixed with 58 grams of Sb₂O₁ are digested with about 50 c c of water till all the Sb₂O₃ goes into solution. The solution is filtered and concentrated gently on the water bath. On cooling, crystals of ammonium antimonyl tartrate separate. Yield=11 grams. The salt is purified by repeated crystallization. It contains ½ molecule of water of crystallization and its antimony content=38.58% on theoretical calculation. Found Sb=38.1%

ANILINE ANTIMONYL TARTRATE

It is best prepared by heating two gram-molecular weights of acid aniline tartrate and one gram-molecular weight of Sb_2O_3 in the presence of water

7 5 grams of tartaric acid are dissolved in water 4 7 grams of aniline are added to this and the mixture boiled for a quarter of an hour. The solution is filtered and crystallized yielding 10 grams of acid aniline tartrate 4 9 grams of acid aniline tartrate are digested with 2 9 grams of anilimony trioxide in the presence of water, till all the antimony trioxide goes into solution. The solution is then allowed to crystallize. Yield=5 1 grams

COMPOSITION

Calculated for C_tH₃NH C₁H SbO O₆ Sb=31 75 o

It has been prepared in other ways by previous workers which need not be described here

Purity of the salts used -

- (1) The sodium and potassium antimonyl tartrates were specially prepared for me as chemically pure by Messrs Martindale & Co.
- (2) The ammonium urea and aniline antimonyl tartrates were prepared and purified in my laboratory by repeated crystallization

The Antimony Content of the Salts Used as Estimated by Actual Calculation

- (1) Ammontum antimonyl tartrate NH_1 C_1H_1 SbO O_6 1H_2O Sb=38 1%
- (2) Urea antimonyl Iartrate CO(NH) (C₁H₃ SbO O₆) ^{1}HO Sh=37 55%
- (3) Aniline antimonyl tartrate C₄H₃NH C₄H₃ SbO O Sb=31 75%
 - (4) Tartar emetic KC4H,SbO O6 HO Sb=36 1.0
- (5) Sodium antimonyl tartrate NaC₁H₁SbO O₆ 23H O Sh=34 1% *

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Lethal doses -

In the following tables and the subsequent portions of this paper the abbreviations used are explained as follows —

- (1) M L D, the minimum lethal dose, ie, the minimum dose in grams per kilo of body weight which killed all the animals used
- (2) Maj L D, the majority lethal dose, ie, the dose in grams per kilo of body weight which killed 66 per cent only of animals used
- (3) M T D, the maximum tolerated dose, ie, the maximum dose in grams per kilo of body weight which was tolerated by all the animals used
- (4) Maj T D, the majority tolerated dose, ie, dose in grams per kilo of body weight which was tolerated by only 66 per cent of the animals used
- (5) T R toxic range, i e, the range between the minimum lethal dose and the maximum tolerated dose

EXPERIMENTS ON GUINEA-PIGS

Method of Administration and Measurement of Doses

The toxicity experiments on guinea-pigs with the above compounds will be first described in the present paper. The drugs were administered intramuscularly, the injections being given in the outer part of the thigh. The strength of the solution was two per cent in distilled water. In all these experiments, each time the solution was freshly prepared and

of condensation of one molecule of $C_8H_9O_4N$ SbNa and two molecules of $C_8H_{10}O_4N$ Sb, 2 H_2O The percentage of antimony present in the compound prepared in our laboratory corresponds to the formula $C_8H_9O_4N$ SbNa, the exact analogue of atoxyl without any water of crystallization Further observations on this subject will be made later on

The difference in the antimony contents of some of the antimonyl tartrates as estimated in my laboratory, from those quoted above, is due to their containing different molecules of water of crystallization. It is a well-known fact that the water of crystallization may vary in an antimonyl tartrate. See Watt's Dictionary of Chemistry [Editor]

an old or stock solution was never used The smaller doses were always measured by means of a tuberculin syringe graduated in hundreds of a cubic centimetre

TABLE 1

Lethal Effects Obtained from the Administration of a 2 per cent Solution of Ammonium Antimonyl Tartrate to Guinea pigs by Intramuscular Injection

Dengmpr kil of body wght	Numb fau ea pgued	Numb dd	Rm ke
06	(6	MLD
055	4	3	
05	4	3	
045	6	4	M;LD
035	6	2	Матр
03	6	N1	мтр

TABLE II

Lethal Effects Obtained from the Administration of a 2 per cent Solution of Urea Antimonyl Tartrate to Guinea pigs by Intramuscular Injection

D ng mp kl fb dy w ght	Numb of gu	1 mb dd	R m 1
055	4	4	MLD
05	2		
045	4	2	
04	6	3	
035	5	2	
03	3	1	MıTD
025	4	N1	MTD
02	1	N1	

TABLE III

Lethal Effects Obtained from the Administration of a 2 per cent Solution of Potossium Antimonyl Tartiate to Guinea-pigs by Intramuscular Injection

Dose in gram per kilo of body weight	Number of gumea- pigs used	Number died	Remarks
055	4	4	MLD,
05	3	2	Maj L D
045	4	2	
04	8	5	
035	6	3	
025	6	3	
02	6	2	Maj T D
015	3	Nıl	мтр

TABLE IV

Lethal Effects Obtained from the Administration of a 2 per cent Solution of Sodium Antimonyl Taitrate to Guinea-pigs by Intramusculai Injection

Dose in gram per Lilo of body weight	Number of guinea- pigs used	Number died	Remarks
055	4	4	MLD
05	3	2	Maj L D
045	4	2	
04	7	5	
035	6	3	
025	4	2	
02	5	1	
015	3	Nil	МТО

PLATE XXIII

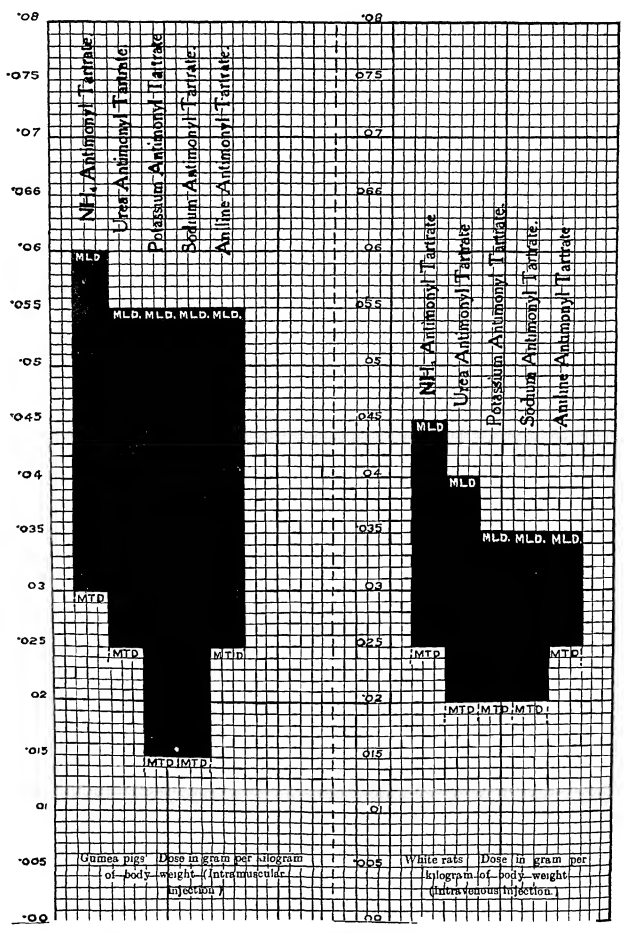


Diagram of a curve formed showing the values obtained for the MLD and MTD of various antimonyl tartrates (Vide page 27)

TABLE V

Leihal Effects Obtained from the Administration of a 2 per cent Solution of Aniline Aniimonyl Tarirate to Guinea pigs by Inframuscular Injection

Digmpr klofbody wght	Number of gu n p gs used	Nmbrded	Rm ke
55	4	4	MLD
05	4	3	
042	6	4	MilD
04	5	3	
03	4	,	
025	7	NI	мто
	1	•	1

Represented graphically the values obtained for the minimum lethal doses and the maximum tolerated doses of the various antimonyl tartrates for guinea pigs will form a curve shown in the accompanying diagram (Plate XXIII)

THE INFLUENCE OF THE BASE OF AN ANTIMONYL TARTRATE UPON ITS TOXICITY

The toxicity of a drug when administered to the same species of animals as determined from its minimum lethal dose is inversely proportional to its minimum lethal dose From this the toxicity of the antimonyl tartrates and of their antimony content can be expressed as follows—

GUINEA PIGS

A (1) Toxicity of ammonium antimonyl tartrate
$$= \frac{K}{06}$$
(2) Do of urea antimonyl tartrate
$$= \frac{K}{055}$$
(3) Do of potassium antimonyl tartrate
$$= \frac{K}{055}$$

(4) Toxicity of sodium antimonyl tartrate
$$= \frac{K}{055}$$

(5) Do of aniline antimonyl tartrate
$$\frac{K}{055}$$

B (1) Toxicity of the antimony content of ammonium antimonyl tartrate

$$=\frac{K}{06\times38\text{ i}}$$

(2) Do do of urea antimonyl tartrate
$$= \frac{K}{055 \times 37.55}$$

(3) Do do. of potassium antimonyl tartiate
$$= \frac{K}{055 \times 36.1}$$

(4) Do do of sodium antimonyl tartrate
$$= \frac{K}{055 \times 34 \text{ l}}$$

(5) Do do of aniline antimonyl tartiate
$$= \frac{K}{055 \times 31.75}.$$

If T (NH₄), T(Urea), T(K), T(Na) and T(Aniline) represent the toxicity of the above tartiates respectively, we then have —

$$\frac{T(NH_4)}{T(Urea)} = \frac{T(NH_4)}{T(K)} = \frac{T(NH_4)}{T(Na)} = \frac{T(NH_4)}{T(Aniline)} = \frac{55}{60} \text{ or } \frac{11}{12}$$

If T Sb (NH₄), T Sb (Urea), T Sb (K), T Sb (Aniline) and T Sb (Na) represent the toxicity of the antimony content of the above tartrates we have —

$$\frac{T \text{ Sb(NH_4)}}{T(\text{Urea})} = \frac{41}{46} \qquad \frac{T \text{ Sb(NH_4)}}{T \text{ Sb(K)}} = \frac{40}{46} \qquad \frac{T \text{ Sb(NH_1)}}{T \text{ Sb(Na)}} = \frac{38}{46}$$

$$\frac{T \text{ Sb(NH_4)}}{I \text{ Sb(Anline)}} = \frac{35}{46}$$

Therefore in the case of the guinea pigs, ammonium antimonyl tattrate is the least toxic then comes the urea salt then the sodium and potassium salts—which are equally toxic and then the amiline salt

The maximum tolerating capacity of the same species of animals for a drug is directly proportional to its maximum tolerated do e

We thus have -

(I) Maximum tolerating capacity of guinea pigs treated with ammonium antimonyl tartrate

				=K' × 03
(2)	Do	do	do	Urea antimonyl tartrate =K' × 025
(3)	Do	do	do	Potassium antimonyl tartiate =K' × 015
(4)	Do	do	do	Sodium antimonyl tartrate =K' × 015
(5)	Do	do	do	Aniline antimonyl tartrate = K' × 025

From this we conclude that of all the antimonyl tartrates used in the case of the guinea pigs their maximum tolerating capacity is with ammonium antimonyl tartrate

EXPERIMENTS ON WHITE RATS

Method of Administration and Measurement of Doses

In the case of white rats the drugs were administered intravenously the injection being given into one of the prominent veins of the tail. The strength of the solution was one per cent in distilled water. Whenever the injection was given the time taken in injecting a given volume of the solution was always the same being at the rate of ½ c c per minute. The injections were given by means of a tuberculin syringe and the solutions were always freshly made.

TABLE VI

Lethal Effects Obtained from the Administration of a 1 per cent Solution of Ammonium Antimonyl Tartrate to White Rats by Intravenous Injection

Dose per kılo	Number of rats used	Number died	Remarks
045 grm 04 ,, 035 ,, 03 ,, 025 ,,	4 6 6 8 4	4 4 4 2 N:1	M L D Maj L D M T D

TABLE VII

Lethal Effects Obtained from the Administration of a 1 per cent Solution of Urea Antimonyl Tartrate to White Rats by Intravenous Injection

Dose per kılo	Number of rats used	Number died	Remarks
04 grm 035 ,, 03 ,, 025 ,, 02 ,,	5 7 2 3 4	5 5 1 1 Nil	M L D Mej L D Maj T D M T D

TABLE VIII

Lethal Effects Obtained from the Administration of a 1 per cent Solution of Potassium Antimonyl Tartiate to White Rats by Intravenous Injection

Dose per kilo	Number of rats used	Number died	Remarks
04 grm 035 ,, 03 , 025 ,,	7 7 7 7 5	7 7 4 3 Nul	MLD

ļ

TABLE IX

Lethal Effects Obtained from the Administration of a 1 per cent Solution of Sodium Antimonyl Tartrate to White Rats by Intravenous Injection

D p kil	Numbrofrt u d	Numb d d	Rmak
04 g m	3	3	
035	10	10	MLD
03	6	4	MILD
025	7	4	}
U2	6	N1	MTD

TABLE X

Lethal Effects Obtained from the Administration of a 1 per cent Solution of Aniline Antimonyl Tartrate to White Rats by Intravenous Injection

D p k lo	Numbsf tud	V mp 1 g g	R m k
04 g m	5	5	
035	6	4	MLD
03	4	2	}
025	5	N1	MTD

Represented graphically the values obtained for the minimum lethal doses and the maximum tolerated doses of the various antimonyl tartrates for rats will form a curve shown in the accompanying diagram (Plate XXIII)

It will be seen from the above tables that the toxic range (T $\,R$) is not so great in the case of white rats as in the case of guinea pigs and the difference in the toxicity of the various antimonyl tartrates is more marked in the case of white rats

Some Aryl Pentavalent Antimony Compounds p-Amino-phenyl Stibinic Acid and some of its Derivatives—their Toxicity

Before proceeding to the study of the arcmatic antimonial compounds dealt with in the present paper, I give here a brief summary of the principles of chemotherapy which have been followed in preparing them

- (1) Pnenyl stibine acid should be more toxe than p-amino-phenyl stibine acid, just as phenyl arsenic acid is more poisonous than p-amino-phenyl arsenic acid.
- (2) The sodium salt of p-amino-phenyl st bin'c acid is the antimony analogue of atoxyl which is of marked therapeutic value in protozoal diseases. I have observed that the urea salt is more stable and less toxic than the sodium salt
- (5) With the idea of reducing the toxicity of p-aminophenyl stibinic acid and its salts, acyl substitution compounds may be prepared by the introduction of various acidic radicles into the amino group of p-amino-phenyl stibinic acid to form secondary amines. Those that have been prepared are described as follows:—
 - (a) Acetyl-p-amino-phenyl stibinic acid and its sodium salt. The latter is identical with "Stibenyl" of Allen and Hanbury.
 - (b) Benzene-sulphonyl-p-amino-phenyl stibinic acid and its sodium salt. This latter is allied to Hectine of Mouneyrat.
 - (c) N-phenyl-glycine-amide-p-stibinic acid and its sodium salt. The above-mentioned acid is allied to N-phenyl-glycine-amide-p-arsenic acid of Jacobs and Heidelberger which has been found by Pearce and Brown to have low toxicity but marked therapeutic properties in experimental trypanosomiasis.

 (Journal of Experimental Medicine, 1919)

- (d) The urcthano derivative of pamino phenyl stibinic acid has been found useful in fowl spirillosis This compound may be described as carbethoxy pamino phenyl stibinic acid
- (4) Allyl thio carbamino p stibanilic acid allied to allyl thiocarbamino p ar anilic acid has been prepared with theidea of having the therapeutic action of allyl and stibinic compounds without the toxic character of the latter
- (5) To reduce the toxicity of the compound phenol p stibinic acid, carboxy methylene group may be introduced into this compound to replace the H of the OH pre ent in the para position of the phenolic compound giving ri e to carboxy methylene oxyphenyl 4 stibinic acid. The corresponding arsenic compound posses as such trypanocidal power that it can cure animals infected with highly resistant strains of trypanosomes.
 - (6) That the introduction of acidic groups into the mole cule of a compound may markedly diminish the therapeutic value of a drug has been taken into consideration
 - (7) When an antimonial compound has to be used for therapeutic purposes
 - $\begin{array}{c} \underline{C} \\ \underline{T} \end{array} \ \ _{1} \ \underline{e} \\ \begin{array}{c} \underline{C} \\ \underline{T} \\ \underline{C} \\ \underline{T} \\ \underline{C} \\ \underline{T} \\ \underline{C} \\$

It will be seen later that (7) can only be determined indirectly in the case of kala azar. In this paper the term effective dose of a drug will be used to denote a dose per day by which the best effect appears to be obtained in the treatment of kala azar when that dose is given for a sufficient length of time. It is impossible to produce therapia sterilisans magna in kala azar with a single dose of any antimonial preparation known up to the present time.

We have now to investigate how far some of the above principles of chemotherapy which are based on theoretical considerations, are borne out by actual experiments

The starting material in the preparation of the new aromatic antimonial compounds dealt with in the present paper is acetyl-p-aminophenyl stibinic acid. The sodium, salt of this compound is sometimes known as stib-acetin. Kala-azar and other forms of leishmaniasis have been successfully treated by its administration (G. Caronia, 1916, Pediatria Also Kharina-Marinucii). More recently, the same compound was used by Manson-Bahr in the treatment of kala-azar under the name of "Stibenyl." (Lancet, Vol. II, 1920)

The successful use of this compound in the treatment of leishmaniasis naturally leads one to attempt to prepare derivatives of this compound allied to those of p-arsanilic acid. The present paper contains a description of some new pentavalent antimony analogues of such derivatives of p-arsanilic as have been found to be of definite therapeutic value

Acetyl-p-amino-phenyl-stibinic acid, CH₁ CO NH C₆H₁ SbO (OH)₂ was prepared in my laboratory by the action of sodium antimonite upon diazo-solution in a way somewhat analogous to Bart's reaction. By diluting the sodium antimonite solution it was found that the yield was greater than that obtained by following the method described in Morgan's work on Organic Compounds of Arsenic and Antimony, which is the method of Von Heyden By the latter method, the preparation is difficult and the yields are low (Percy May) The percentage of antimony in C₈H₉O₄ N Sb Na, the sodium salt of the above acid is 36 8 By actual calculation it was found to be 36 1

This compound yields on hydrolysis 4-amino-phenyl-stibinic acid (Von Heyden D R P, 270, 488). The sodium salt of this acid is the antimony analogue of atoxyl and has been described in the German patent (Von Heyden D R P, 254, 421)

For the sake of simplicity I have called this sodium salt NH C_cH_iSbO OH stibamine from its analogy to the

corresponding salt of arsenic which is also known as ars amine (Journal of Tropical Medicine and Hygiene August 15 1921) Stibamine as prepared in my laboratory is an amorphous powder fairly soluble in water. Its solution helps its decomposition. The solution should be freshly prepared before use.

Composition -

Calculated for C₆H O₃ NSb Na Sb=42 25% N=4 93% Found Sb=42 10%, N=4 88%

(1) Urea stibamine CO (NH)₂ H₂ N C H₄ SbO (OH)

This is carbamide salt of p amino phenyl stibinic acid [This compound was prepared with the idea that the urea compound could be used intramuscularly possessing anæs thetic properties as quinine urea — Editor]

EXPERIMENTAL.

2 3 grams of p amino phenyl stibinic acid suspended in water are treated with solid urea until a clear solution is obtained on slight heating. The solution is then concentrated on the water bath. To the concentrated solution absolute alcohol is added in excess when a precipitate forms. The mixture is heated for a few minutes to dissolve any excess of urea. The precipitate is then filtered and thoroughly washed with absolute alcohol to dissolve the last traces of uncombined urea. It is then dired on porous plate. The yield is about 1.5 grams. I propose to call this compound urea stibamine.

The salt is fairly soluble in water and is amorphous

Composition —

Calculated for $C_7 H_{12} O_1 N_3 Sb$, Sb = 37 26%, N = 13 04%

Found Sb = 36.95%, N = 12.52%

The to a and therapeutic properties of this compound will be described in the present paper

(2) Benzene-sulphonyl-p-aniino-phenyl-stibinic acid

C₆ H₇ SO₂ NH C₆ H₁ Sb O₃H₂

The sodium salt of this compound is the antimony analogue of Hectine of Mouneyrat, which is sodium-benzene-sulphonyl-p-amino-phenyl arsinate. This latter compound is of marked reputed value in spirochæte infection.

EXPERIMENTAL

Benzene-sulphonyl-p-amino-phenyl-stibinic acid

5 gram of stibamine is dissolved in 2 c c of $\frac{N}{1}$ sodium hydroxide and treated with 5 gram of benzene-sulphonyl chloride. The mixture is warmed on water bath at 60°C and shaken from time to time. The alkali is replenished as soon as it is found to be exhausted. After an hour and a half, the reaction is found to be complete. The solution is filtered and conc. HCl is added to it drop by drop until it is distinctly acid. The sulphonyl compound is precipitated and is then filtered. For purification, the precipitate is suspended in water and carefully dissolved in $\frac{N}{1}$ sodium hydroxide and again precipitated by hydrochloric acid. The process is repeated three times and the precipitate is filtered and carefully washed with water and dried on a porous plate. The yield is 33 gram.

ć

The sodium salt is a fairly stable compound and is freely soluble in water—It has not yet been obtained in a crystalline form—I propose to call it stib heetine

Composition -

Calculated for C_1 H_1 O NS Sb, Sb=29 85% N=3 5% Found Sb=29 3% N=4 06%

(3) Urethane Derivative of p amino phenyl stibinic acid (carbethoxy p amino phenyl stibinic acid)

EXPERIMENTAL

2 9 grams of p amino phenyl stibinate of sodium and 6 gram of 35 per cent caustic soda solution are dissolved in 10 c c of water. The mixture is treated with 1 3 grams of ethyl chlorocarbonate and 1 2 grams of 35 per cent caustic soda solution and stirred. After about half an hour the mixture is filtered. From the filtrate urethano derivative is precipitated with dilute hydrochloric acid. The precipitate is purified by dissolving it in caustic soda solution and then precipitating again with dilute hydrochloric acid.

Composition --

Calculated for C, H₁ O₅ N Sb Sb=35 92%, N=4 19 N Found Sb=35 74% N=3 96%

(4) Carboxy methylene oxyphenyl 4 stibinic acid

COOH CH, O C.H. ShO, H

EXPERIMENTAL

It is prepared by adding an alkaline solution of mono chloracetic acid to the solution of sodium phenol p stibinate

Preparation of p hydroxy phenyl stibinic acid —The diazo solution obtained from a mixture of 2 2 grams of p amino phenol 3 grams of sulphuric acid in 20 c c of

water, and 1'4 grams of sodium nitrite is added with stirring, to a sodium antimonite solution. The latter is obtained by mixing a solution of antimony trichloride, prepared by dissolving 2-88 grams of antimony trioxide in 12 c.c. of hydrochloric acid (D=1-123) and an aqueous solution of sodium hydroxide (12 grams) in 120 c.c. water. When the decomposition is complete, the excess of sodium hydroxide is almost neutralised with dilute sulphuric acid. The mixture is saturated with carbon dioxide and filtered repeatedly to remove any traces of antimony trioxide. p-hydroxy-phenyl-stibinate of sodium is then precipitated by saturating the solution with sodium chloride. p-hydroxy-phenyllstibinic acid is precipitated from the solution of the latter in water by dilute sulphuric acid. It is then filtered and dried on porous plate.

Preparation of carboxy-methylene-oxyphenyl-4-stibinic acid—It is prepared by adding successively monochloracetic acid (1 88 grams) in 3 c c of water and 4 grams of 35 per cent caustic soda solution to the solution of 2 63 grams of p-hydroxy-phenyl-stibinic acid and 4 gram of caustic soda in 5 c c of water. The mixture is heated at 60°C, for about three hours. When cooled, the mixture is carefully acidified with hydrochloric acid. The precipitated acid is purified by dissolving in sodium hydroxide solution and precipitating again with dilute hydrochloric acid.

(5) N-phenyl-glycine-amide-p-stibinic acid NH₂ OC H₂ C NH C₆ H₁ SbO (OH)₂

I have tried to investigate whether amino-phenyl-stibinic acid possesses the property of giving rise to compounds of the following type RHN CO CH_2 NH C_6H_1 SbO (OH)₂ which are similar in constitution to those prepared by Jacobs and Heidelberger from p-arsanilic acid (Journal of American Chemical Society, 1919) Of these glycine compounds of antimony, I have prepared N-phenyl-

glycine amide of stibinic acid which is allied to N phenyl glycine amide of arsenic acid of the above authors. This latter compound has given very remarkable results in the treatment of experimental trypanosomiasis and spirochæte infection in the hands of Pearce and Brown (Journal of Experimental Mcdicine 1919). It is therefore expected that the corresponding antimony compound prepared in my laboratory should exhibit similar results in the treatment of leishmaniasis. Its toxicity and therapeutic properties have not yet been studied by me

EXPERIMENTAL.

N (phenyl 4 stibinic acid) glycinc amide or N phenyl glycinc amide p stibinic acid

8 gram of stibamine is dissolved in 4 c c of $\frac{N}{1}$ sodium

hydroxide solution After adding 74 gram of chloracet amide the mixture is warmed on water, both under a reflux conden er for about two hours. During warming a reddish brown precipitate is gradually formed and settles at the bottom, the flask being shaken from time to time. After the operation the crude product is allowed to cool 14 cc of concentrated hydrochloric acid is added to the cold mixture to hold any unchanged stibamine in solution During this treatment the portion of the amido glycine compound which was retained in solution by the alkali is precipitated substance is then filtered off and carefully washed with cold For purification it is suspended in sufficient, water to form a thin paste and carefully treated by stirring with sodium hydroxide solution until the acid is dissolved. It is filtered from the undissolved product and is then treated with a little excess of dilute acetic acid whereupon the substance separates as a white precipitate After filtering and washing thoroughly it is quickly dried on a porous plate and kept in a sealed tube The yield is 2 gram The acid is purified by its repeated solution in alkali and precipitation by acetic acid

Sodium Salt — The pure acid is suspended in enough water to form a thick paste and carefully treated with 25 per cent sodium hydroxide solution, until completely dissolved and the solution reacts neutral to litmus. Two volumes of alcohol are then added, the pure sodium salt quickly separating as a white powder. After filtering and washing with 85 per cent alcohol it is quickly dried on a porous plate.

The acid is sparingly soluble in cold water—It dissolves more easily in hot water—The sodium salt is freely soluble in water—It has not yet been obtained in a crystalline form and is less soluble in water than the corresponding arsenic compound—I propose to call this compound stib-glycine-amide

Composition —

Calculated for $C_8 H_{10} O_1 N_2 Sb Na$, Sb = 35 19%, N = 8 2%

Found Sb=3541%, N=79%

(6) Allyl-thio-carbamino-p-amino-phenyl-stibinic acid C₃ H₅ NH CS NH C₆ H₁ SbO (OH)₂

The above compound is prepared by treating stib-amine with allylthiocarbamide in methyl alcohol

EXPERIMENTAL

2 gram stibamine is dissolved in 3 c c of methyl alcohol and to the mixture 08 grm oleum sinapis (containing 90 per cent allylthiocarbamide) is added. The mixture is kept at ordinary temperature for 24 hours, and then filtered. The filtrate is diluted with a little water and treated with a few drops of concentrated hydrochloric acid, which precipitates the allylthiocarbamino derivative. The crude product is filtered and washed with water and dried in a desiccator. The dried substance is finally washed with ether, to free it from oil. The compound obtained is a yellowish white

amorphous powder soluble in sodium hydroxide solution but not soluble in sodium carbonate The yield is 2 gram

Composition -

Calculated for C_{10} , O_3 H_{11} , N S Sb Sb = 33.2% N = 7.7%

Found Sb=33 4% N=7 8%

In the present paper the toxicity and therapeutic proper ties of urea stibamine will be described. The toxicity of some of the other aryl antimonial compounds dealt with in the present paper will also be described here.

Toxicity Experiments with Phenyl Stibinate of Sodium Stibamine Urea Stibamine, etc

Method of administration —The drugs were administered into guinea pigs intramuscularly the injections being given in the outer part of the thigh. The strength of the solution was 2 per cent in distilled water. In all these experiments, each time the solution was freshly prepared and an old solution was never used.

TABLE XI

Lethal Effects Obtained from the Administration of a 2 per cent Solution of Phenyl Stibinate of Sodium to Guinea pigs by Intramuscular Injection

D p kıl	Nmb of gu pg	ď	Numbrd d	Rmk
2 g m			2	1
1	2		2	1
05	3		3	
6,40	4		N i	{

TABLE XII

Lethal Effects Produced from the Administration of a 2 per cent Solution of acetyl-p-amino-phenyl-stibinate of Sodium to Guinea-pigs by Intramuscular Injection (Stibenyl)

Dosc per l'ilo	Number of guines pigs used	Number died	Remarke
7 grm	3	3	MLD
6	5	3	
5 ,,	1	2	
45 ,,	1	2	design
4 ,,	6	1	† 1
35 ,	2	Nil	MID

TABLE XIII

Lethal Effects Produced from the Administration of a 2 per cent Solution of Urea Stibamine to Guinea-pigs by Intiamuscular Injection

Dose per kilo	Number of guinea pigs used	Number dicd	Remarks
7 grm	4	1	MLD
65 ,,	3	2	Maj L D
6 ,,	4	2	
5 ,,	2	1	
45 ,,	4	1	
4 ,,	4	l l	
35 ,,	4	Nıl	MTD

TABLE XIV

Lethal Effects Produced from the Administration of a 2 per cent Solution of Stibamine to Guinea pigs by Intramuscular Injection

D s p l lo	N mb of gun pg u d	Numbrd d	Rmk
J g m	4	4	MLD
45	4	3	
4	4	3	1
35	4	2	
3	8	2	
2	6	NI	MTD

TABLE XV

Lethal Effects Produced from the Administration of 2 per cent Solution of Stib hectine to Guinea pigs by Intramuscular Injection (Compound made in my Laboratory)

Doep ki	Nmb fgnapgud	Nambe d d	Rmrk
6 g m	4	4	
5	4	4	MLD
4	3		Ì
3	3	2)
2	2	1	}

TABLE XVI

Lethal Effects Produced from the Administration of a 2 per cent Solution of Stib-hectine to Guinea-pigs by Intramuscular Injection (Compound supplied by Chemisch Fabrik von Heyden)

Dose per kilo	Number of guinea pigs used	Number died	Remarks
5 grm	3	3	
4 ,,	5	5	MLD
3 ,,	4	3	
25 ,,		1	

Symptoms of poisoning after intramuscular injection of aryl antimonials into guinea-pigs—These symptoms are, generally speaking, similar to those following toxic doses of the antimonyl tartrates. The pathological changes in the organs after toxic doses of the aryl antimonials are also similar to those obtained after administration of toxic doses of antimonyl tartrates. In one case in which the optic nerve was examined, I did not find such degenerative changes in the optic nerve as have been observed to have followed the use of the aryl arsonates.

Having determined the toxicity of stibamine and urea stibamine, I give a summary of their physical and chemical properties

Physical and Chemical Properties of Stibamine and Urea-stibamine

Properties of stibamine —

- (1) Stibamine is a brown amorphous powder soluble in water, the solution being of a reddish-yellow colour
- (2) The solution of stibamine is easily decomposed, giving rise to a precipitate containing antimony, in the presence of an acid or alkali

(3) The filtrate after separation of the above precipitate also contains antimony

Properties of urea stibamine -

- (1) It is a brown amorphous powder like stibamine and is soluble in water giving rise to a reddish solution. It is in soluble in alcohol
- (2) Unlike stibamine its solution is not so easily decomposed by boiling for a few minutes. Its solution can be sterilized by boiling
- (3) It is more stable than stibamine when kept in solution
- (4) It is an additive compound of urea and liberates N_2 when treated with a solution of sodium hypobromite

The therapeutic value of ammonium antimonyl tartrate and usea stibamine —

Having proved that ammonium antimonyl tartrate is the least toxic of the five antimonyl tartrates investigated in the present paper. I now pass on to describe its effects when administered to man for therapeutic purposes. Only a few clinical cases will be described here. It is however beyond the scope of the present paper to give the comparative very allue of the various antimonyl tartrates in the treatment of kala azar.

TREATMENT OF KALA AZAR WITH INTRAVENOUS INJECTION OF AMMONIUM ANTIMONYL TARTRATE

(1) Patient K æt 30 was admitted into hospital suffer ing from kala azar. The spleen extended 5¹ below the costal margin. On spleen puncture L D bodies were found At the time of admission the body weight was 5 stone Patient was treated with intravenous injections of ammonium antimonyl tartrate twice a week the doses being increased from 2 c c to 8 c c of a 2 per cent solution. Altogether 6-7678

16 injections were given As a result of the treatment, the fever of the patient completely stopped, the spleen could not be felt below the costal margin and on spleen puncture no L. D. bodies could be found after the 16th injection. Patient increased one stone in weight during the treatment

Result of Blood Examination —

- (1) R.B'C —2,300,000, W.B C —2,000, Hb—32 per cent on 15-6-1921 before treatment
- (2) R B C —4,300,000, W B C —7,000, Hb—55 per cent on 12-9-1921 after treatment.
- (3) Patient N, æt 16, was admitted into hospital suffering from kala-azar The spleen extended 4" below the costal margin. On spleen puncture many L D bodies were found. At the time of admission, the body weight was 3 stone. Patient was treated with intravenous injections of ammonium antimonyl tartrate twice a week, the doses being increased from 1 c c to 6 c c of a 2 per cent solution. Altogether 25 injections were given. As a result of treatment, the fever of the patient completely stopped, the spleen could not be felt below the costal arch after the 20th injection, and at the time of discharge no L D bodies could be found on spleen puncture. Patient increased 1 stone in weight during the treatment.

Result of Blood Examination —

- (1) R B C —3,200,000, W B C —2,600 Hb—44 per cent on 3-6-1921 before treatment
- (2) R B C —4,500,000, W B C—7,000, Hb—60 per cent on 1-6-1921 after treatment
- (3) Patient A, æt 21, was admitted into hospital suffering from kala-azar The spleen extended 7" below the costal margin. At the time of admission the body weight was 6 stone. Patient was treated with intravenous injections of ammonium antimonyl tartrate twice a week, the doses being

increased from 3 cc to 9 cc of a 2 per cent solution. Altogether 14 injections were given. As a result of treatment, the fever subsided, the spleen could just be felt below the costal margin after the 15th injection and at the time of discharge no L. D. bodies could be found on spleen puncture. Patient increased, I stone in weight during the treatment.

Result of blood examination -

- (1) R B C -2 900 000 W B C -1 200 Hb-42 per cent on 30 6 1921 before treatment
- (2) R B C -- 4 200 000 W B C -- 11 400 Hb-- 55 per cent on 5 11 1921 after treatment

Each of the above cases appeared to be cured the will be seen that the highest dose given up to now was 9 c c of a 2 per cent solution. Symptoms of vomiting and purging were not great after these injections. A series of cases which could not bear treatment with tartar emetic on account of severe reactions such as high fever vomiting and purging are now being treated with ammonium antimonyl tartrate with less marked reactions. The intramuscular injection of the compound is painful and may give rise to local reaction which is not so marked as in the case of tartar emetic.

TREATMENT OF KALA AZAR WITH INTRAVENOUS INJECTION OF UREA STIBAMINE

In the following cases of kala azar the effects of intra venous injection of urea stibamine are briefly recorded

(1) Name—Manu æt 10 years L D bodies found on spleen puncture Dose= 15 gram given twice a week

Effect of treatment —

R.B C	W.B C	Hb
4,200,000	3,200	50 per cent on admission
3,100,000	4,800	40 ,, after 5 injections
3,400,000	4,800	48 ,, ,, 16 ,,
3,200,000	10,400	46 ,, ,, 20 ,,

Spleen reduced from 3½" to 1½" below the costal arch No L D bodies found after 20 injections Patient free from fever for one month

(2) Tofu, æt 30 years L D bodies found on spleen puncture before treatment

Dose =
$$\begin{cases} (1) & 10 & c & c - 2 \text{ injections } (0.2 & \text{gram}) \\ (2) & 12 \\ 2 & c & c - 13 \\ 3 & 15 & c & c - 3 \end{cases}$$
, (0.25 gram)

Injections given twice a week

Effect of treatment —

RBC	WBC	НЬ
2,900,000	1,800	38 per cent on admission.
2,800,000	4,200	38 ,, after 3 injections
3,200,000	5,200	42 ,, ,, 14 ,,
4,700,000	10,400	52 ,, ,, 18 ,,

Spleen reduced from $2\frac{1}{2}$ " to *nil* beneath costal arch. Body weight increased from 5 st $2\frac{1}{2}$ lb to 6 st 3 lb No L D bodies found on spleen puncture after 18 injections Patient free from fever for one month.

(3) Abdul, æt 30 years Spleen 6" below the costal arch in the left nipple line and 2" away from mid-line to the right side L D bodies found on spleen puncture before treatment

Dose = 2 gram at each injection Injections given twice

Effect of treatment -

RBC	WBC	НЬ
1 900 000	008 1	28 per cent on admission
3 000 000	2 600	44 , after 7 injections
3 900 000	3 600	50 13

Spleen slightly felt below the costal arch and body weight increased from 6 st 6 lb to 7 st 4 lb Patient discharged at his own request Patient free from fever for one month

(4) Anath Bondhu, at 14 years L D bodies found on spleen puncture before treatment

Dose = 2 gram twice a week

Effect of treatment -

RBC	WBC	HЬ
2 600 000	1 200	38 per cent on admission
2 800 000	4 200	42 after 5 injections
3 800 000	000 8	50 , 12

Spleen reduced from 6 to almost nil below the costal arch Body weight increased from 4 st 21 lb to 5 st 6 lb Patient free from fever for one month

(5) Abdul act 12 years Spleen 4½ below the costal arch L D bodies found on spleen puncture before treatment

Dose = 15 gram twice a week

Effect of treatment -

-	- ur by realin			
	RBC	WBC	H)
	2 900 000	1 400	44 per cent on	admission
	3 700 000	2 400	48 af	ter 10 injections
	4 200 000	2 400	50	14
	3 900 000	5 200	42	16

Patient absconded from hospital

(6) Mosafar, æt 30 years L D. bodies found on spleen puncture before treatment

Doses — 1st $2\frac{1}{2}$ c c , 2nd 5 c c., 3rd 10 c c and the last 6 injections $12\frac{1}{2}$ c c of a 2 per cent solution. Injections given twice a week

Effect of treatment —

RBC	WBC	Hb
2,400,000	1,600	32 per cent on admission.
2,400,000	7,000	36 ,, after 9 injections.

Patient absconded from hospital

(7) Horoz, æt 10 years L D bodies found on spleen-puncture before treatment Spleen extended 5½" below the costal arch before treatment.

Dose = 05 to 15 gram every alternate day

Effect of treatment —

RBC	WBC,	Нь
2,300,000	2,000	36 per cent before treatment
3,400,000	3,800	40 ,, after 3 injections
4 200,000	13,800	46 , ,, 16 ,, and
		same after 27 injections

Spleen just felt below costal arch. Patient free from fever for one month

No L D bodies found on spleen puncture after 20 injections.

(8) Abdul, æt 25. L D bodies found on spleen puncture before treatment

Dose = 25 gram twice a week

Effect of treatment -

RBC	WBC	Hb		
3 000,000	2 400	40 per cent on admission		
4 300 000	7 000	50 after 7 injections		

Spleen reduced from 7 to almost nil below the costal arch Body weight increased from 6 st to 7 st 4 lb

Patient left hospital before treatment was completed

REMARKS

In the present paper the toxicity of some of the antimonyl tartrates and some of new aromatic antimonials has been described

In his observation on the Treatment of Oriental Sore Greig has come to the following conclusions with regard to the use of tartar emetic in the treatment of the disease

It is not desirable to exceed 12 to 13 cc (1 per cent solution) at one time as toxic symptoms become more marked above this limit. Hence we see that the $\frac{C}{7}$ dose ratio of

antimonium tartaratum is not very satisfactory the organo and parasito tropic properties are not in the correct proportion

The ideal drug for the destruction of Leishmania tropica in the tissues has still to be sought (Indian Journal of Medical Research October 1917) The same also holds good in the use of the drug in the treatment of kala azar. The average minimum effective dose of tartar emetic in the case of an adult man in the treatment of kala azar may be taken as 6 c c of a 2 per cent solution (= 12 gram). The average minimum effective dose of urea stibamine used for the same purpose is 25 gram. If C and C¹ denote these

minimum effective doses respectively we have $\frac{C}{C^{2}} = \frac{12}{25}$ or

³ approximately

From the toxicity experiments described in the present paper it will be seen that the maximum tolerated doses per kilo of body weight in the case of the guinea-pig are '015 gram of tartar emetic and 35 gram of urea stibamine. If T and T' represent these tolerated doses we have —

$$\frac{T}{T'} = \frac{015}{35}$$
 or $\frac{1}{23}$ nearly

Though it does not necessarily follow that the minimum tolerated dose for the human being can be reckoned weight for weight by rule of three with mathematical accuracy from observations on the guinea-pig, still it is evident from the above figures that urea stibamine is a much safer antimonial for use in the treatment of kala-azar than tartar emetic. The fact also holds good in the case of the other antimonyl tartrates. The effective dose of urea stibamine in the treatment of kala-azar is ? this the tolerated dose for the guinea-pig, while in the case of tartar emetic, it is 8 times the tolerated dose for the same animal

Conclusions

- (1) After the administration of a toxic dose of an antimonyl tartrate, the pathological changes are most markedly seen in the lungs, the kidneys, the liver, pituitary and adrenals. These consist chiefly of hæmorrhages into the substance of these organs and destruction of their cellular elements. Similar pathological changes are also observed after toxic doses of the aromatic antimonial compounds
- (2) Ammonium antimonyl tartrate is the least toxic of all the antimonyl tartrates used
- (3) The toxicity of the antimony content of an antimonyl tartrate is least marked in the case of the ammonium salt.

•

- (4) The presence of N in the basic radicle of an antimonyl tartrate diminishes the toxicity of some of them
- (5) Ammonium antimonyl tartrate is of marked thera peutic value in the treatment of kala azar
- (6) The low toxicity of ammonium antimonyl tartrate and its high antimony content lead to the conclusion that of all the antimonyl tartrates dealt with in the present paper ammonium antimonyl tartrate is the best for use in the treatment of hala azar
- (7) A series of new organic aromatic antimonials have been discovered the preparations of which have been described in the body of the paper
- (8) The toxicity of the following aromatic antimonials has been estimated in the case of the guinea pig (1) Phenyl stibinic acid (2) Acetyl p amino phenyl stibinic acid (3) Stibamine (4) Urea stibamine (5) Stib hectine
- (9) The acetyl derivative of p amino phenyl stibinic acid is less toxic than stibamine
 - (10) Urea stibamine is less toxic than stibamine
- (11) Urea stibamine has been found useful in the treatment of kala azar
- (12) Urea stibamine is a much safer antimonial for use in the treatment of kala azar than tartar emetic or other antimonyl tartrates
- (13) Symptoms such as vomiting and purging are much less marked after intravenous injection of an effective dose (= 25 gram) of urea stibamine than that of tartar emetic or sodium antimonyl tartrate (= 12 gram)
- I am deeply indebted to my staff Mr Niharranjan Chatterjee M Sc Mr Saradacharan Chaudhury M Sc Dr Pramathanath Ghose M B and Sub Assistant Surgeon Bibhutibhushan Maity L S M F for helping me in carrying

on my researches My grateful thanks are also due to Mr Parimal Sen, M.Sc, for the drawings and sections of all the organs described in this paper.

[N B.—The paper of Fargher and Gray on the Chemotherapy of Antimony, which was published after my paper was sent to the Secretary, Indian Research Fund Association, last December, will be discussed in a subsequent communication.]



First case of Dermal Leishmanoid
(Reproduced from a paper of the author published in the Indian Medical Gazette, Vol LVII, No. 4,
April, 1922)

CHEMOTHERAPY OF ANTIMONIAL COMPOUNDS IN KALA AZAR INFECTION

PART II

DERMAL LEISHMANOID

[Recei ed f r Publi at on Jun 5 1922]

Under the name of Derinal Leishmanoid I described in the April 1922 issue of the Indian Medical Gazette a form of dermal leishmaniasis which developed in a case of kala azar cured by antimonial treatment. Since the publication of this paper. Major Knowles IMS has made a series of inoculations and cultural experiments. which are described below.

I am indebted to the Editor Indian Medical Gazette for permission to reproduce here a drawing showing the eruptions on the upper part of the patient's body (Plate LXXIII) A drawing from the scrapings from one of the papules is also appended herewith showing the presence of Leishmania donovani which seem to be mostly extra corpuscular in the smear A few have been found inside leucocytes and endothelial cells (Plate LXXIV Fig 1)

INOCULATION AND CULTURAL EXPERIMENTS

A series of experiments have been performed to determine if flagellates developed from the Leishman Donovan bodies which had apparently been modified in their virulence by a course of antimonial treatment and also to determine if they could infect monkeys by giving rise to a local or general disease

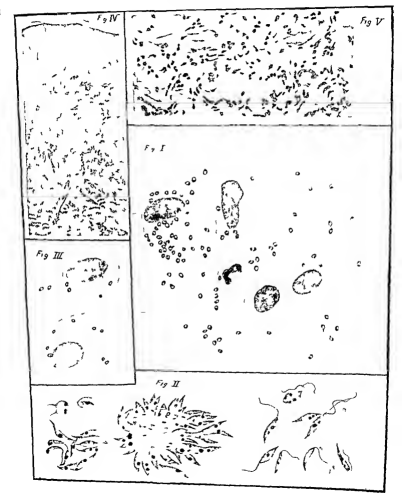
The following are the notes on the cultural and inoculation experiments very kindly made for me by Major Knowles, I.M.S., Protozoologist, Calcutta School of Tropical Medicine:—

- (1) One of the nodules of the right arm was pricked and the serum which cozed out was inoculated into NNN medium and incubated at 22°C Flagellated bodies were found after 12 days, and these were indistinguishable from those of Leishmania donovani (Plate LXXIV, Fig. 2)
- (2) The culture from the peripheral blood of the patient and smears from the same gave negative results Examination of the splenic blood—negative
- (3) A monkey (*M thesus*) was inoculated in both eyebrows by embedding bits of granulomatous nodules into pockets cut in them. After a month-and-a-half, marked granulomatous growths were observed over the sites of inoculation in both eyebrows. (See Diagram.) Also small secondary nodules were observed at the outer and inner canthuses of the eyes. One of the nodules at the original site of inoculation was incised and smears made from it—a fair number of Leishman-Donovan bodies were present, most of which were intra-corpuscular and a few extra-corpuscular and free. (Plate LXXIV, Fig. 3.)
- (4) No ulceration has yet been observed over the nodules after two-and-a-half months, the raw surface left after incising one of the nodules having healed up
- (5) Blood examination and culture from the peripheral blood of the monkey were negative a month-and-a half after inoculation
- (6) The smears from the liver of the monkey and cultures from the same organ on NNN medium gave negative results a month-and-a-half after inoculation

- Fig 1, Plate LXXIV—scrapings from one of the papules, showing the presence of L D bodies which are mostly free forms and extracapsular, a few are inside leucocyte and endothelial cells (Vide also p 53)
- Fig 2, Plate LXXIV—scrapings showing flagellated bodies found after 12 days which were indistinguishable from those of L D bodies
- Fig. 3, Plate LXXIV, shows a fair number of L. D. bodies in smears from one of the incised nodules taken from the original site of inoculation made in the monley
- Fig 4, Plate LXXIV—section of a papule in the skin of the patient showing round-celled infiltration with fibroblasts and thinning af the epidermis
- Fig 5 is the same as Fig 4 showing the presence of a network of newly formed capillaties and thickening of capillary walls

[R p t df m th Ind a Jou n Lof M d 1R r h Vol X No 4 Ap | 1923]

PLATE LXXIV





[Reprint d from th 1 don] in lof Medic I Re es h 1 of X No 4 Ap 1 1923] FLATE LXXIV (b)



Phig phof m ky hwag m kd g nimitu gowth tih t f nocult n nboth y b ws d n a m nth nd h lib f by emb dd ng bt f grulm t n d! into pock t cut n th m with a c nd ryn dul tih outrnd nn c nth f th ey



HISTO PATHOLOGY OF THE GRANULOMA

The normal structure of the chorium is replaced by granulation tissue consisting mainly of large cells which are arranged in columns and between them young fibroblasts and fine capillaries of the granulation tissue are seen and there in the large capillaries the endothelium is hyper trophied and the wall is thickened In some places there is a very marked thickening of the wall of the capillaries almost leading to their obliteration. On the wall of the thickened capillaries and in their endothelial lining are seen Leishman Donovan bodies The skin over the papillomat ous nodules is moderately pigmented and stratum cornium is very thin. The papillæ of the chorium are much less prominent than the normal ones There is no surface ulceration No Leishman Donovan bodies are seen in the epidermis On the whole the pathological changes in the skin are very similar to those recently described by Cornwall in a non ulcerated oriental ore (Plate LXXIV Figs 4 and 5)

The Leishman Donovan bodies are best seen in smears from the scrapings from the granulomatous nodules. In these smears they appear mostly as free parasites. Here and there on careful examination they are found also inside large mononuclear leucocytes and very rarely inside the polynuclears. Besides they are found inside other large cells which are perhaps endothelial cells. (Plate LXXIV Fig. 1.)

After the publication of the case the patient underwent a course of antimonial treatment alternated with intravenous injection of salvarsan and seemed to improve somewhat But he left Calcutta before the completion of his treat ment

Leishman Donovan bodies have been occasionally discovered in the papular eruptions and scrapings from

ulcers in the skin in cases of kala-azar (Christophers) The findings in the above case of dermal leishmanoid, however, differentiate it from such cases, as, so far as can be made out, the present case is purely a local *Lcishmania* infection of the skin in a patient otherwise cured of kala-azar.

OBSERVATIONS

- (1) The granulomatous nodules of the skin contain Leishmania donovani, most of which are extra-cellular and some intra-cellular—the reverse of what occurs in oriental sore
- (2) The culture from the serum from the granulomatous nodules produced flagellate forms indistinguishable from those of Leishmania donovani
- (3) An inoculated monkey developed granulomatous nodules containing Leishmania donovani which are mostly intra-cellular but some are extra-cellular, just the reverse of what was found in the patient and more resembling the findings in oriental sore
- (4) The disease in the case of the monkey is still a local disease. But no ulceration has yet been observed in the nodules of the inoculated monkey.
- (5) The patient's peripheral blood and splenic blood gave negative results
- (6) The pathological changes in the skin are very similar to those observed in oriental sore

The facts, established by the cultural and inoculation experiments described in the present paper, that the disease is purely a lesion of the skin, that no Leishmania could be cultivated from the peripheral blood, that the examination of the patient's splenic blood was negative, and that in the successfully inoculated monkey no Leishmania could be cultivated from the peripheral blood and the liver, all appear to support my view that the disease was due to a modified

virus of Leishmania donovani. The histological changes in the skin resemble those of a non ulcerated oriental sore with this difference that in the present disease the Leishmania are mostly extra corpuscular while in oriental sore they are mostly intra corpuscular. The reverse condition has however been found in the monkey. These facts lead me to conclude that the modified virus of Leishmania donovani brought about by antimonial treatment resembles that of Leīshmania tropica the causative agent of oriental sore.

CONCLUSIONS

The case is therefore one of cutaneous leishmaniasis due to Leishmania donovani and proves for the first time that these parasites can sometimes produce only skin manifestations in man without visceral lesions. This condition has followed antimonial treatment of kala azar

My grateful thanks are due to Major Knowles 1 M S for the invaluable help he has given me in making the cultural and inoculation experiments. To my assistant Mr Parimal Bikas Sen M Sc. I am greatly indebted for the drawing of the Plates for my paper.

CHEMOTHERAPY OF ANTIMONIAL COMPOUNDS IN KALA-AZAR INFECTION

PART III

FURTHER OBSERVATIONS ON THE TOXICITY OF ANTIMONIAL COMPOUNDS—DELAYED ANTIMONY POISONING

[Received for Publication, December 19, 1922]

(A)

FURTHER OBSERVATION ON THE TOXICITY OF ANTIMONIAL COMPOUNDS

EXPERIMENTS ON GUINEA-PIGS

The same methods of administration and measurement of doses were followed as in my previous paper. These will not therefore be described again.

(1) Lithium antimonyl taitrate

Method of preparation —

EXPERIMENTAL

37 grm of lithium carbonate is slowly added to a watery solution of 1.5 grms of tartaric acid. The mixture is heated till all the CO₂ is expelled. To the solution 1.44 grms of Sb₂O₃ are added and the mixture gently heated till all the Sb₂O₃ dissolves. The solution is filtered and concentrated and to the concentrated solution is added four times its volume of absolute alcohol. Lithium antimonyl tartrate is precipitated as granular crystals.

Yield—24 grams
Calculated for LiC₁H₄O Sb Sb=412 per cent
Found Sb=406 per cent

Lethal effects obtained from the administration of a two per cent solution of lithium antimonyl tartrate into guinea pigs by intramuscular injection —

TABLE I

Dose in gr m per kilo	No of animal used	Number d ed
045	4	4
64	6	, 6
03	4	3
02	4	IN

Toxicity of antimony content = $\frac{k}{04 \times 406}$ or $\frac{k}{1624}$

(2) Calcium antimonyl tartrate

Method of preparation -

EXPERIMENTAL

56 grm of anhydrous calcium oxide is dissolved in a watery solution of 3 grms of tartaric acid. To the mixture 2 88 grms of 5b O₁ are slowly added while it is heated on the sand bath. When all the antimony trioxide has gone into solution the solution is filtered concentrated and allowed to crystallize. It is purified by recrystallization from water.

Yield-5 grams

Heated at 110°C found loss of weight = 148 per cent Calculated for CaCsH₅O₅₆Sb 64HO weight of water of crystallization = 161 per cent Calculated for the above hydrated salt, Sb=33 1 per cent. Found Sb=33 0 per cent

Lethal effects produced from the administration of a two per cent solution of calcium antimonyl tartrate into guinea-pigs by intramuscular injection —

Dose in gram per kilo	No of animals used	Number died
055	2	2
05	4	3
045	4	3
02	4	0

TABLE II

Toxicity of antimony content = $\frac{K}{055 \times 33}$ or $\frac{K}{1815}$

(3) Strontium antimonyl taitrate

Method of preparation —

EXPERIMENTAL

To a watery solution of 15 grms of tartaric acid is slowly added 73 grms of strontium carbonate. The mixture is then heated till all the CO₂ is expelled. The solution is then heated on the sand bath and during the process 144 grms of Sb₂O₃ are slowly added to it and the heating is continued till all the Sb₂O₃ dissolves. The solution is then filtered, concentrated and allowed to crystallize. It is purified by crystallization from water.

Yield—3 grams

Calculated for SrC₅H₅O₁₄Sb₂, Sb=3666 per cent Found Sb=366 per cent. Lethal effects produced from the administration of a two per cent solution of strontium antimonyl tartrate into guinea pigs by intramuscular injection —

|--|

No fram Israed	A mt rd d
3	3
4	3
7	6
,	1
2	1
2	1
	3 4 7 2 2 2

Toxicity of antimony content =
$$\frac{k}{0.55 \times 36.6}$$
 or $\frac{k}{2.01}$

(4) Ethyl antimonyl tartrate

Method of preparation -

Ethyl antimonyl tartrate was prepared in solution by Prof Collie by heating freshly precipitated antimony trioxide with acid ethyl tartrate to about 150°C in a sealed tube [Pio Roy Sco 82 B (1910)—252) The latter can be prepared after the method described by Guerin Varry—A 1837 22 238 (vide Sudborough s Practical Organic Chemistry) An other method is that of J Bongault [J Pharm Chim 1906 (VI) 23 465 69]

The best method of preparing ethyl antimonyl tartrate is by the action of ethyl iochde on silver antimonyl tartrate in presence of absolute alcohol

EXPERIMENTAL

Silver antimonyl tartrate is prepared by adding a solution of silver nitrate (2 6 grams) to a concentrated solution of

tartar emetic (5 grams). This yields white granular crystals of silver antimonyl tartrate. The precipitated silver salt is filtered, washed with cold water and then dried in air.

41 grms of silver antimonyl tartrate dried at 105°C. are mixed in a conical flask, with about 20 c.c of absolute alcohol and excess of ethyl iodide. The mixture is gently heated on the water bath for about five hours under reflux. After the completion of the reaction a yellow undissolved mass, containing ethyl antimonyl tartrate and silver iodide, is obtained. The mixture is then filtered and washed with absolute alcohol. The residue is then extracted with a small quantity of cold water. The ethyl antimonyl tartrate is obtained from the solution by gently evaporating it on water bath

Yield—About 2 grams.

It is a white shining powder coluble in water. The solution is distinctly acid to litmus and can be sterilized by gentle boiling without decomposition

Calculated for $C_2H_5C_4H_4O_7Sb$, Sb=38.3 per cent. Found Sb=37.8 per cent.

Lethal effects obtained from the administration of a one per cent solution of ethyl antimonyl tartrate into guinea-pigs by intramuscular injection —

 Dose in gram per kilo
 No of animals used
 Number died

 05
 2
 2

 045
 6
 6

 04
 3
 1

 03
 4
 3

 02
 4
 2

TABLE IV

Toxicity of the antimony content = $\frac{K}{045 \times 37.8}$ or $\frac{K}{1.701}$

Method of preparation of antimonyl tartrates of quinine and eigebonine

(5) Quinine antiquonyl tartrate

EXPERIMENTAL

1 9 grms of barium antimonyl tartrate are dissolved in 400 c c of water and to the solution are added 2 8 grms of quinine sulphate. The whole is vigorously shaken from time to time for 24 hours after which it is filtered when the filtrate is found to be free from sulphate and barium. The filtrate is now kept in the vacuum desiccator for drying. It is very spannigly soluble in water.

Calculated for $C_{24}H_{\nu_0}O_{\nu_0}N$ Sb Sb=19.7 per cent Found Sb=19.3 per cent In attempting to prepare quinine antimonyl tartrate by boiling Sb₂O₃ with acid quinine tartrate a portion of the quinine is converted into the toxic quinotoxine

(6) Cinchonine antimonyl tartrate

EXPERIMENTAL

1 18 grms of cinchonine base is heated on the water bath containing 6 grm of tartaric acid until all the cinchonine goes into solution. To the above solution 8 grm of Sb₂O₃ is added and the mixture heated on the water bath until all the antimony trioxide goes into solution. The solution is filtered concentrated and allowed to crystallize. The crystals appear to be globular. They are purified by recrystallization from water.

Calculated for C_3H O_8N Sb Sb=20 7 per cent Found Sb = 20 6 per cent The compound can also be prepared in the same way as quinine antimonyl tartrate

EXPERIMENTAL

I grm of barium antimonyl tartrate is dissolved in water and to this is added a saturated solution of 1 l grams of cinchonine sulphate. The mixture is slightly heated on the water bath in order to complete the reaction. It is then filtered and the filtrate kept over sulphuric acid, and allowed to crystallize, when globules of crystals of cinchonine antimonyl tartrate appear. They are purified by recrystallization.

Calculated for $C_{23}H_{27}O_8N_2$ Sb, Sb=20 7 per cent Found in the dried substance Sb=20 7 per cent

Lethal effects obtained from the administration of a solution of quinine antimonyl tartrate into guinea-pigs by intramuscular injection —

TABLE V

Dose in gram per kilo	No of guinea pigs used	Number died
15	3	3
125	5	4
I	6	5
05	2	Nıl

Toxicity of the antimony content =
$$\frac{K}{15 \times 19.7}$$
 or $\frac{K}{2.955}$

Lethal effects obtained from the administration of a solution of cinchonine antimonyl tartrate into guinea-pigs by intramuscular injection —

TABLE VI

Dose í grm pe kil	No figures pig used	N mber d d
15	,	2
125	2	1
1	4	4
09	2	2
08	3	,
-0	2	1
re	4	3
05	1	2
		l

Toxicity of the antimony content = $\frac{k}{15 \times 201}$ or $\frac{k}{3015}$

(7) Narcoline antimonyl tartrate

Method of preparation -

EXPERIMENTAL

4 12 grms of narcotine are digested with a watery solution of 1.5 grms of tartaric acid till all the narcotine dissolves. The solution is concentrated and two grms of antimony trioxide are added and the mixture heated on water bath for some time. The mixture is then diluted with water and filtered. The filtrate is concentrated on the water bath. After cooling needle shaped crystals of narcotine antimonyl tartrate are obtained which are purified by recrystallization. It has no water of crystallization.

Yield—5 grams nearly Calculated for C H₂O₁₁NSb Sb=17 2 per cent Found Sb=16 8 per cent Lethal effects obtained from the administration of a solution of narcotine antimonyl tartrate into guinea-pigs by intramuscular injection —

TABLE VII

Dose in gram per l'ilo	No of guinea pigs used	Number died
ì	2	2
085	4	1
08	4	3
07	4	2
06	2	1
055	2	0
05	2	0

Toxicity of the antimony content = $\frac{K}{085 \times 16.8}$ or $\frac{K}{1.4}$

EXPERIMENTS ON RATS

Lethal effects obtained from the administration of a one per cent solution of lithium antimonyl tartrate into rats by intravenous injection —

TABLE VIII

Dose in gram per kilo	No of animals used	Number died
03	4	4
025	2	2
02	5	5
015	6	5
01	- 1	0

Lethal effects obtained from the administration of a one per cent solution of calcium antimonyl tartrate into rats by intravenous injection —

TABLE IX

Do in gram per k lo	, 1	No of a matus d	Numbrd d
03		3	3
025	1	3	1
02	1	1	ſ
015		2	. 1
-01		2	, 0

Lethal effects obtained from the administration of n one per cent solution of strontium antimonyl tartrate into rats by intravenous injection —

Tyble X

Dose in g m per k lo	No of mtued	Nmbrded
03	2	2
025	1	1
70	2	1
015	1	, 1
- 01	1	0

Lethal effects produced from the administration of a one per cent solution of ethyl antimonyl tartrate into rats by intravenous injection —

TABLE XI

D i g am per k l	N famlused	Numb dd
03	4	4
025	6 /	5
02	3	2
015	1	0
1		

Lethal effects produced from the administration of a solution of quinine antimonyl tartrate into rats by intravenous injection —

TABLE XII

Dose in gram per kilo	No of animals used	Number died
1	4	4
09	2	1
08	4	3
07	4	1
06	2	1

Lethal effects produced from the administration of a solution of cinchonine antimonyl tartrate into rats by intravenous injection.—

TABLE XIII

Dose in gram per kilo	No of animals used	Number died
08	1	1
07	2	2
06	4	3
'05	2	0

TOXICITY OF OLD SOLUTIONS OF TARTAR EMETIC
AND OF OLD SAMPLES OF STIBENYL

It has been found by many observers that toxic symptoms may follow intravenous injections of old solutions of tartar emetic. This leads one to investigate whether an old solution is more toxic to guinea-pigs and rats than fresh solutions. I give here the results of toxicity experiments with such solutions.

A 2 per cent solution of tartar emetic was kept in an Erlenmeyer flask and the white precipitate that is frequently formed in old solutions was allowed to increase After three weeks the solution was made up to the original volume and injected after sterilization along with the precipitate into guinea pigs and the following results were obtained

Lethal effects obtained from the indministration of an old (three weeks) 2 per cent solution of turtar emetic into guinea pigs by intramuscular injection —

TABLE XIV

Deigmperkilo	No of nim 1 sd	N mbrd d
04 03 02 015	3 2 2 6	3 1 1 0

Therefore M L D with guinea pigs = 04 grm per kilo while in the case of fresh solution M L D = 055 grm per kilo lt is thus evident that solutions of tartar emetic become more and more toxic in course of time

In the case of white rats similar results were obtained as will be seen from the following tables

Lethal effects obtained from the administration of an old (three weeks) 2 per cent solution of tartar emetic into white rats by intramuscular injection —

TABLE XV

D ng mpe klo	No of nm ls d	Numb d d
05 045 04 035 03 075	2 9 4 3 3	2 2 9 4 3 2

Lethal effects obtained from the administration of a fresh two per cent solution of tartar emetic into white rats by intramuscular injection —

TABLE XVI

Dose in gram pei kilo	No of animals used	Number died
04	4	4
035	5	3
03	4	3
025	2 ′	1
02	2	0

Therefore M L D. in the case of white rats with old solutions of tartar emetic = 03 gram per kilo of body weight given intramuscularly, while with fresh solutions it is '04 per kilo of body weight. Investigations are in progress to determine whether the increase in toxicity is due to any change in the optical activity of the solution

Toxicity of Old Samples of Stibenyl

The first samples of stibenyl that were supplied to me towards the end of 1920 were tested by me after a year and a half and the following results were obtained

Lethal effects obtained from the administration of a solution of old (nearly a year and a half) sample of stibenyl into guinea-pigs by intramuscular injection —

TABLE XVII

Dose in gram per kilo	No of animals used	Number died
6	2	2
4	4	4
35	2	1
25	1	1

Therefore the M L D of old samples of stibenyl is 4 gram per kilo of body weight while that of fresh samples is 7 gram per kilo of body weight. It therefore follows that stibenyl kept in powder form becomes more and more toxic in course of time lift may be stated here that no difference in solubility could be observed in these old samples of stibenyl the substance in each case quickly going into solution.

TOXICITY OF ANTIMONYL MALATES

The two antimonyl malates that I have so far investigated are (I) Ammonium antimonyl malate (2) Sodium antimonyl malate. In the process of purification of these salts they are found to crystallize with a molecule of sodium or ammonium hydrogen malate with formation of double salts. Of these salts. I have found that the ammonium salt is more stable while the sodium salt is easily decomposed in solution on boiling.

EXPERIMENTAL

Sodium antimonyl malate —About 4 grms of Sb O₀ are digested with an aqueous solution of 6.2 grms of acid sodium malate till no further antimony trioxide is dissolved. The operation is conducted on the water bath under reflux for about an hour when the reaction is complete. The solution is filtered concentrated and then allowed to crystallize from water.

Yield-6 grams

Ammonium antimonyl malate —About 4 grms of Sb O_3 are digested with 4.5 grms of acid ammonium malate in the same way as the above till no more of the Sb O_3 goes into solution. The process is then conducted in the same way as above. The double salt is purified by recrystallization from water.

Yield—5 grams

It contains five molecules of water of crystallization Calculated for $C_8H_{27}O_{16}N_2Sb$, Sb=22.7 per cent Found Sb=22.61 per cent

Lethal effects obtained from the administration of a two per cent solution of the double salt of ammonium antimonyl malate and acid ammonium malate into guinea-pigs by intramuscular injection —

Dose in gram per kilo	No of animals used	Number died
125	4	4
09	5	5
08	4	2
075	2	1
07	4	0

TABLE XVIII

Toxicity of the antimony content of the salt = $\frac{K}{09 \times 2261}$ or $\frac{K}{203}$

Double Salt of Sodium Antimonyl Malate and Acid Sodium Malate

The toxicity of this salt cannot be determined with accuracy, as it decomposes on boiling during the process of sterilization of its solution

TOXICITY OF Sb₂O₃ DISSOLVED IN GLYCERINE

Lethal effects obtained from the administration of a two per cent solution of Sb_2O_3 in glycerine into guinea pigs by intramuscular injection —

7	LADE	e.	X	v

eng mpklo	No of manufu d	Numbrd d
045	4	4
05	4	4
03	4	4
025	4	1
02	4	0

Antimony content of Sb₂O₃=83 per cent

Toxicity of the antimony content =
$$\frac{k}{03 \times 83}$$
 or $\frac{k}{249}$

TOXICITY OF Sb O, DISSOLVED IN TARTARIC ACID

 $Sb\ O_s$ gives a series of acids when it combines with tartaric acid. These need not be enumerated here

In the following experiments Sb O₃ was dissolved in the minimum quantity of tartaric acid and the strength of the solution used was in terms of Sb O₃ dissolved in tartaric acid

Lethal effects obtained from the administration of a two per cent solution of SbO in tartaric acid into guinea pigs by intramuscular injection —

TABLE XX

D ng mp kl	No of mlud	N mb dd
035	6	6
03	4	4
025	6	3
02	4	1
015	4	0

Toxicity of the antimony content = $\frac{k}{249}$

Yield-5 grams.

It contains five molecules of water of crystallization Calculated for $C_8H_{27}O_{16}N_2Sb$, Sb=22.7 per cent Found Sb=22.61 per cent

Lethal effects obtained from the administration of a two per cent solution of the double salt of ammonium antimonyl malate and acid ammonium malate into guinea-pigs by intramuscular injection —

Dose in gram per lilo	No of animals used	Number died
125	4	4
09	5	5
08	4	2
075	2	1
07	4	0

TABLE XVIII

Toxicity of the antimony content of the salt = $\frac{K}{09 \times 2261}$ or $\frac{K}{203}$

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TOXICITY OF Sb₂O₃ DISSOLVED IN GLYCERINE

Lethal effects obtained from the administration of a two per cent solution of Sb₂O₃ in glycerine into guinea pigs by intramuscular injection —

T	_	v	v

D ng mp kl	No of an m is s d	Numb d d
045	4	4
05	4	4
03	4	4
075	4	1
02	4	0

Antimony content of Sb O₃=83 per cent

Toxicity of the antimony content =
$$\frac{K}{03 \times 83}$$
 or $\frac{K}{249}$

TOXICITY OF Sb O₃ DISSOLVED IN TARTARIC ACID

Sb O_2 gives a series of acids when it combines with tartaric acid. These need not be enumerated here

In the following experiments Sb O_3 was dissolved in the minimum quantity of tartaric acid and the strength of the solution used was in terms of Sb O_2 dissolved in tartaric acid

Lethal effects obtained from the administration of a two per cent solution of ${\rm Sb_2O}$ in tartaric acid into guinea pigs by intramuscular injection —

TABLE XX

D g mp klo	Noofanmlud	V mp q q
035	6	6
03	4	4
025	6	3
02	4	1
015	4	0

Toxicity of the antimony content = $\frac{k}{249}$

COMPARISON OF THE TOXICITY OF THE VARIOUS ANTIMONIALS SO FAR INVESTIGATED AND OF THE RESULTS OBTAINED WITH THOSE OF OTHER OBSERVERS

Making a summary of the various antimonials so far investigated by me and calculating on the basis that their toxicity is inversely proportional to their minimum lethal doses, we find that their toxicities in the case of guineapigs × 10 are as follows —

TOXICITY OF THE ANTIMONY CONTENT OF ANTIMONYL TARTRATES AND MALATES IN THE CASE OF GUINEA-PIGS

	•	
Quinine antimonyl tartrate	= 1 30 Cinchonine antimonyl tartrate	- 1 30
Sb2O1 dissolved in glycerine	= 1 25 , Sb_O3 dissolved in taitaire acid	= 1 25
Ammonium antimonyl tartrate	= 1 23 Urea antimons! tartrate	= 1 21
Strontrum antimonyl tartrate	= 1 20 Ammonium antimonyl malate	= 1 20
Potassium antimonyl tartrate	= 1/20 Sodium antimony I tartrate	= 1119
Calcium antimonyl tartrate	= 1/18 Anilme antimonyl tartrate	= 1 17
Ethyl antimonyl tartrate	= 1 17 Lithium antimonyl tartrate	= 1 16
Narcotine antimony I tartrate	= 1[14 '	

The experiments of Farghar and Gray—I have not however been able to confirm the observations of these two workers that the toxicity of the antimony content of quinine antimonyl tartrate is only one-fifth that of tartar emetic, though I agree with them that its toxicity is less than that of tartar emetic I confirm their observations that quinine antimonyl tartrate on boiling with antimony trioxide is converted into the more toxic quino-toxine antimonyl tartrate. I have not been able to confirm their, as well as Rogers', conclusions that the sodium salt is less toxic than the potassium salt. I have confirmed Plimmer and Thompson's observations that the lithium salt is more toxic than the sodium or potassium salt.

I have been able to prepare the antimony analogue of atoxyl without its water of crystallization. Farghar and Gray seem to have obtained the sodium salt of a polymenzed derivative of acetyl p amino phenyl stibinic acid which they could not completely free from the admixture of sodium chloride. By determining the molecular weight of the compound by the freezing point method after dissolving it in distilled water it was found that the molecular weight was 266.6 which more nearly corresponds to the constitution of the compound given by me than to that given by Farghar and Gray. My observations on the minimum lethal doses with the antimonyl salts so far investigated on guinea pigs give higher figures than those obtained by Farghar and Gray in the case of mice.

It would appear from the observations of the above workers that the least toxic antimonyl tartrate to be used in the case of kala azar should be quinine antimonyl tartrate The effective dose of tartar emetic in the case of kala azar is generally 5 c c of a two par cent solution (= I grm or 15 grains) According to their observations 5 times the effective dose of tartar emetic can be used in the case of quinine antimonyl tartrate. This will be 75 grains The amount of quinine base present in 7.5 grains of quinine antimonyl tartrate is nearly 4 grains. Apart from any other effect of quinine the intravenous injection of 4 or 5 grains of quinine in a concentrated solution given rapidly may sometimes lead to a dangerous fall of blood pressure especially in a weak and debilitated kala azar patient and therefore should not be advocated I have discussed the effect of the fall of blood pressure after intravenous injection of guinne elsewhere

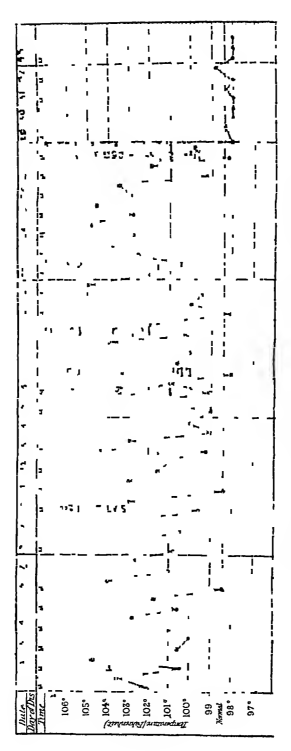
Amongst others who have worked on the toxicity of antimony compounds may be mentioned Carl Voegtlin and co workers who experimented with albino rats Korns experimented with rabbits

(B) DELAYED ANTIMONY POISONING

In some rare cases, after intramuscular injection of an antimonyl tartrate or malate into guinea-pigs, I have found that death took place even so late as three weeks or more after one injection. The viscera which showed pathological changes were subsequently subjected to chemical examination and showed the presence of antimony. The organs examined were the liver, the lungs and the kidneys. These cases may be described as cases of delayed antimony poisoning.

The details of post-moitem examination of a number of guinea-pigs that died of delayed antimony poisoning are as follows —

- (1) Serial No 449—Guinea-pig weighing 237 grms was given an intramuscular injection of a two per cent solution of tartar emetic in dose of 04 grm per kilo of body weight Total quantity of tartar emetic injected was 0095 gram. The animal died 22 days after the injection. On post-moitem examination, there were hæmorrhages in both the lungs Liver was fatty and there were patches of necrosis here and there over the surface of the liver. Kidneys were congested. The animal evidently died of antimony poisoning, the viscera showing the presence of antimony.
- (2) Serial No 445—Guinea-pig weighing 455 grms was given an intramuscular injection of a two per cent solution of sodium antimonyl tartrate in dose of 04 grms per kilo of body weight. Total quantity of sodium antimonyl tartrate injected was 018 grm. The animal died 14 days after the injection. On post-mortem examination there were hæmorrhages in the gall bladder. The kidneys were congested. The animal evidently died of antimony poisoning, the viscera showing the presence of antimony.
- (3) Serial No 426 —Guinea-pig weighing 620 grms was given an intramuscular injection of a two per cent



Temperature chart of a kala-azar case treated successfully with urea stibamine

solution of double salt of namonium natimonyl malate and ocid ammonium malate in dose of 5 grm per kilo of body weight. The total quantity of namonium antimonyl malate injected was 262 grm. The animal died three weeks after the injection. On post mortem examination there were hemorrhages in both the lungs. The liver was fatty and the kidneys congested. The animal evidently died of antimony poisoning the viscera showing the presence of ontimony.

- (4) Serial No. 439—Guiner pig weighing 495 grms was given an intramuscular injection of one per cent solution of tartar emetic in dose of 5 grm per kilo of body weight. The total quantity of tartar emetic injected was 025 grm. The naimal died 19 doys after the injection. On post mortem examination there were hemorrhoges in the left lung. There was a entorthal condition of the whole gostro intestinal tract. Liver was fotty and there were patches of necrosis on the surface. kidneys were congested. Traces of ontimony were found in the liver the lungs, the kidney and the intestines. Animal died of ontimony poisoning.
- (5) Serial No 432—Guineo pig weighing 160 grms was given on intramuscular injection of n one per cent solution of tartar emetic in dose of 04 grm per kilo of body weight. Total quantity of tartar emetic injected was 0064 grm. The animal died three weeks after the injection. On post mortem examination there were lizemorrhages in the right lung. The gastro intestinal tract was ulcerated. The liver was fatty and there were patches of necrosis on its surface. The kidneys were congested. On chemical examination of the liver the kidneys the lungs and the intestines there was distinct presence of antimony. The animal died of antimony poisoning.
- (6) Serial No 463—Guinea pig weighing 200 grms was given an intramuscular injection of o one per cent solution of sodium antimonyl tartrate in dose of 045 grm

per kilo of body weight. The total quantity of sodium antimonyl tartrate injected was 009 grm. The animal died 18 days after the injection. On post mortem examination, there were hæmorrhages in the right lung. The liver was fatty and there were patches of necrosis on the surface. The kidneys were congested. There were distinct traces of antimony in the viscera which showed the pathological changes. The animal died of antimony poisoning

(7) Senal No 450.—Guinea-pig weighing 227 grms was given an intramuscular injection of a one per cent solution of tartar emetic in dose of 04 grm per kilo of body weight. The total quantity of potassium antimonyl tartrate injected was 0091 grm. The liver was fatty. The kidneys were pale and somewhat enlarged in size. There was distinct presence of antimony in the liver, the kidneys and the lungs. The animal died of antimony poisoning 26 days after injection.

METHOD OF CHEMICAL EXAMINATION OF VISCERA FOR DETECTION OF ANTIMONY

The organs are cut into small pieces with a pair of scissors. Then these are digested under slow heat with chemically pure HCI (1 in 4), to which a piece of bright copper foil, free from arsenic or antimony, is added. If arsenic or antimony is present, then there is a deposit on the copper. The piece of copper is then washed with alcohol and then with some ether and subsequently made absolutely dry. Then it is put into a hard glass reduction tube and heated for some time. The sublimate obtained on the cold portion of the tube is examined under the microscope, and if antimony is present then an amorphous deposit or sometimes characteristic needle-shaped crystals of Sb₂O₃ are obtained. If the sublimate is sufficient, it can then be dissolved in dilute HCl and H₂S passed into the solution. An orange coloured precipitate shows the presence of Sb in the solution.

Cases of delayed antimony poisoning are of very great clinical importance as they prove that the excretion of the drug may sometimes be very slow after injection of antimonial compounds and some of the cases of sudden death during antimonial treatment may be due to a cumulative action of the drug

REMARKS

- The toxicity of the following antimonial compounds has been worked out in the present paper (I) Lithium antimonyl tartrate (2) Calcium antimonyl tartrate (3) Stroatium antimonal tartrate (4) Ethal antimonal tartrate (2) Quinine antimonyl tartrate (6) Cinchonine antimonyl tartrate (7) Narcotine antimonyl tartrate (8) Ammonium antimonyl malate (9) Sb O, dissolved in glycerine and (10) Sb.O. dissolved in tartaric acid. By comparing their toxicities it was found that ouinine antimonyl tartrate is one of the least toxic antimonial salts. But for reasons that guinting is likely to bring about a dangerous fall of blood pressure after an intravenous injection its antimonyl tartrate cannot be recommended for use in place of ammonium sodium or potassium antimonyl tartrates Sb O₄ dissolved in glycerine or tartaric acid comes next in order of low toxicity but for obvious reasons the solution cannot be recommended for use intravenously SbO, dissolved in tartaric acid is the basis of the author's hyp racid antimonyl tartrate which has given satisfactory results in the treatment of kalanzar when used intramuscularly SbO, dissolved in glycerine is the basis of Martindale s injectio antimonii oxidi
 - 2 Old solutions of tartar emetic and old samples of stibenyl have been found to be more toxic than fresh ones
 - 3 Cases of death in guinea pigs three weeks or so after one injection of an antimonial salt have been met with showing definite symptoms of antimony poisoning and presence of antimony in the viscera

CHEMOTHERAPY OF ANTIMONIAL COMPOUNDS IN KALA-AZAR INFECTION

[Received for Publication, December 19, 1922]

PART IV

FURTHER OBSERVATIONS ON THE THERAPEUTIC VALUE OF URFA STIBAMINE.

New series of cases

(1) Patient, named Puran, æt 12 years, was admitted into hospital with history of double rise of temperature Spleen extended 7 inches below the costal margin in the left nipple line. Body weight—3 st. 2 lbs. Patient was treated with intravenous injection of urea stibamine twice a week. During the treatment patient had frequent attacks of dysentery. Fever stopped after five injections. Altogether 20 injections were given, each dose being 1 gram and the duration of treatment four months. Patient was under observation for nearly a month after treatment was stopped during which there was no fever and at the time of discharge, the spleen could not be felt below the costal margin nor could L. D. bodies be found on spleen puncture. Body weight—3 st. 8 lbs.

Blood count	Spleen puncture	Remarks
R B C -2,100,000, W B C1,800, Hb-32% R B C -4,200,000, W B C -6 200, Hb-54% R B C -4,200 000, W B C -7,800, Hb-52%	Positive Negative Negative	Before treatment 120 days after commencement of treatment Total amount injected —2 grams in 20 injections after which treatment was discontinued 26 days after treatment was discontinued when patient was discharged

ŧ

(2) Patient named Jatin at 25 years was admitted into hospital with double rise of temperature Spleen—74 inches below the costal arch Body weight—5 st 4 lbs Patient was treated with intravenous injection of urea stiba mine twice a week in doses of 5 c c to 73 c c of n 2 per cent solution (= 1 to 15 grm) As a result of treatment the fever stopped after three injections Patient was under observation for nearly a month after the injections were stopped and at the time of discharge spleen could not be felt below the costal margin Body weight—6 stone

Pleod count	Spl n p neture	REMARKS
R B C -2 300 000 W B C -1 400 Hb -36%	Pos ti	Befte tm nt
R B C - 2 903 032 W B C - 2 400 Hb-44 s	-	18 day afte t tm nt w ti n d 6 inject n h ing be n gi n Amount i ject d—8 grm
R B C -3 500 000 W B C -4 200 Hb -48	Ngi	45 dy fir mm cm tof t tm t 12 leet hig beeng: aft wild inc tin w dec tud Am t leet d-17 sim
R B C = 3 800 000 W B C = 5 400 Hb = 57%	Neg t	35 day firt imnt w deted whn the pilit we dehed

(3) Patient, named Bepin was admitted into the hospital for treatment of kala azar with continuous fever Spleen—4 inches below costal arch Body weight—6 st 71 lbs Patient was treated with intravenous injection of urea stiba mine twice a week in doses of 5 c c of a 2 per cent solution (= 1 grm) There was slight rise of temperature with rigor after the first five injections The fever came down to

normal after the sixth injection. The spleen could not be felt below the costal margin, when the treatment was stopped. The patient was under observation for nearly three weeks after the injections were stopped, during which there was no fever and at the time of discharge not D bodies could be found on spleen puncture. Body weight 7 stone

Blood count	Spleen puncture	Remas
7 · ·	-	;
R B C = 3,100 000 W B C = 1,400 Hb=11 ,	Posts c	1 Before treatment
R B C -2,000 000 W B C -2 800, Hb-10		20 days after counsers of the treatment of injections having here are a Around invested 55 pits.
R B C - 3,400 000, W B C - 6,100 Hb - 52	Negative	63 ds, other controlled is a testment to injection have been steers bight the special discontinued. Amount injected—152-min.
R B C -3,400,600, W B C -7,000 Hb-52°,	Negative	24 days after treatment was clickly timed as hen the patient and discharged

(4) Patient, named Abdul, set 20 years, was admitted into the hospital for treatment of kala-azar with fever Spleen—3½ inches below costal arch. Spleen puncture—L D bodies found Body weight—4 st 18 lbs. He was treated with intravenous injection of urea stibamine in doses of 5 c c to 7½ c c of a 2 per cent solution (= 1 grm. to 15 grm.) There was slight nausea without any other reaction after each injection. The spleen could not be felt below the costal margin shortly after the completion of injection. He was 30 days under observation after injections were stopped, during which there was no rise of temperature. At the time of discharge no L. D. bodies could be

determined on spleen puncture Body weight—5 st 2 lbs

Blode nt	Spleen p ct	Remarks
R B C -2 900 000 W B C -t 600 Hb-40%	Pι	B for tr stm nt
R B C -3 400 000 W B C -3 200 Hb-42		30 dys ft omm met f tr tm t 10 injeton h g bee give Amout nj i d—1 g m
R B C -3 400 000 W B C -3 400 Hb- 45%	Ngt	47 dy fir commenc m ni of t im ni 15 i ject on h ing bee g n aft wh h they w e stopp d Am unt i jected-175
R B C -3 400 000 W B C -6 200 Hb-	N. 1	gem 30 day firitm niwa discontaned who pitwis dis- higed

(5) Patient named Prithi Raj act 30 years was admitted into hospital with very high fever and slightly enlarged spleen. At the time of admission the patient was in a drowsy state which lasted for 4 days after admission the spleen rapidly increasing to 31 inches below the costal arch The blood showed no malarial parasites Body weight after he recovered from his drowsiness-6 stone Patient used to get rise of temp-rature with rigor L D bodies found on spleen puncture The patient was treated with intravenous injection of urea stibamine in doses of 5 c.c. to 7½ c c of a 2 per cent solution (= 1 to 15 grm) The fever stopped after three injections Very few L D bodies were found on spleen puncture after six injections Altogether 15 injections were given and after this the spleen could just be felt below the costal arch at the time of dis charge no L D bodies could be found on spleen puncture Duration of treatment-66 days

Błood count	Spleen puncture	Remare 5
R B C -3,603,030 W B C -1,400 Hb-48'.	Po itive	Before treatment
R B C —4 300,000, W B C —6,400,Hb—50°,	Negative	65 days after commencement of treatment altogether 15 injections liaving licen given. I otal amount injected—2 prims
R B C -4,507 000.	Negative 1	40 days after completion of treat ment
W B C -6,800 Hb-52°,	Negative	69 days after treatment was stopped after which patient was discharged from hospital in excellent condition

(6) Patient, named Monglu, æt 18 years, was admitted into hospital suffering from kala-azar. Spleen-4 inches below costal arch Body weight—3 st 4 lbs There was L D bodies were found on spleen slight jaundice puncture Patient was treated with intravenous injection of urea stibamine in doses of 5 cc to 71 cc of a 2 per cent solution (= 1 to 15 grm) Altogether 20 injections were given He did not get much reaction after the injections, except nausea after injection of $7\frac{1}{2}$ cc of a 2 per cent solution As a result of treatment the fever subsided At the time when the injections were discontinued the spleen did not much diminish in size, but no L D bodies could be found on spleen puncture Patient remained in hospital for 60 days after the injections were stopped, and at the time of discharge the spleen completely disappeared below the costal arch and the patient improved considerably in health Body weight-4 stone Duration of treatment-95 days.

Blo de u t	P pleral bl d ult NVN med um	Spl n		REMARKS
R B C -2 500 000 W B C -1 800 Hb-36~	Polt c	Po t	Po t	Bf trat
R B C -4 500 Mb -52 W B C -4 800 Hb -52				46 dy alt omm cm nt ft tm nt 10 ij t h igb ne Ttal mut- 95 g m
R B C _4500 r00 W B C _5 F00 Hb_55%	Ngte	Neg t	Ngt	95 dy oft cmm m t of fr tun mt 20 i jectin h i geb ng siter which fr tm nt w stopp d T t mount-235 gtm
	Ng t	Ngt	Net	35 dy aft emplion of timit
R B C 4 400 000 W B C 5 800 Hb 58	Neg t	Ngt	Ngu	55 dy fte emplin of tre iment Put 1 ft hop t 100 dy ft mplin of t m t

(7) Patient named Romesh aet 14 years was admitted into the hospital suffering from kala azar with fever and dysentery and oedema of the extremities Spleen 6 inches below costal arch LD bodies found on spleen puncture Body weight 3 stone Patient was treated with intravenous injection of urea stibamine in doses of 2½ c c to 5 c c of a 2 per cent solution (= 05 to 1 grm) in spite of dysentery Developed cancrum oris after eight injections which healed up in course of treatment As a

result of treatment the fever subsided and when the treatment was discontinued, spleen extended 2½ inches below the costal arch, but no L D bodies could be found on spleen puncture. Patient remained in hospital for 70 days after the injections were stopped, and at the time of discharge the spleen completely disappeared under the costal arch Body weight—4 st 2 lbs Duration of treatment—90 days

Blood count	Peripheral blood culture NNN medium	Spleen puncture	Spleen blood culture N N N medium	Remari s
R B C2,600,000, W B C1,200, Hb—32% R B C4,100,000, W B C6,600, Hb—52%	Positive	Positive	Positive	Pelore treatment
,				commen ce- ment of treat- ment, 12 injections having been given Total amount —1 15 grms
R B C —3 700,000, W B C —6,400, Hb—52%	Negative	Negative	Negative	90 days after c o m m e n c e- ment of treat ment, 20 injec- tions h a v i n g b e e n g i v e n after which the injections were stopped Total amount — 1 95 grms
	Negative	Negative	Negative	33 days after completion of treatment
R B C —3,600,000, W B C —7,000, Hb—48%	Negative	Negative	Negative	65 days after completion of treatment, after which the patient left hos pital

(8) Patient named Phaninder set 15 years was admitted into the hospital for treatment of kala azar Spleen—41 inches below costal arch Temperature varied from normal to 103°F Bleeding from the gums and nose present Body weight—5 st 4 lbs * No L D bodies found on spleen puncture Patient was treated with intravenous injection of urea stibamine in doses of 5 c c to 7½ c c of a 2 per cent solution (= 1 to 15 grm) No reactions after the injections Altogether 16 injections were given As a result of treatment the fever stopped after 12 injections when the spleen could hardly be felt below the costal arch After 16 injections no L D bodies could be found on spleen puncture Patient is still in hospital 90 days after completion of treatment Body weight—6 stone Duration of treatment—50 days

El deut	Pph I blod ultu NNN mdum	Spl n pun t	Spl n bl d cult NNN md m	Remarks
R B C -2 300 000 W B C -2 600 Hb-36 R B C -3 700 000	Pte	Ngt	Pt	Bf tr t
W B C —3 700 Hb—48	Ngt	Ngı	Pt	16 dy ft omm n m t f t t m nt 6 1 t h ng b n g n Tot l m unt45 g m
W B C -4 200 Hb-44	Ngt	Ngt	Ngt	50 dy ft omm ne mnt ft t mnt f6 nj t n h ng be n g ft wh h t tm nt w t pp d T t 1 m t - 165 g m

The test LDbd ldbjd plpt

Blood count	Peripheral blood culture NNN medium	Spleen puneture	Spleen blood culture N N N medium	Remarks
R B C5,000,000, W B C8,200, Hb50 %	Negative	Negative	Neg itive	120 days after completion of treatment, patient still in hospital His general condition shows remarkable improvement

(9) Patient, named Haren, æt 12 years, was admitted into the hospital with spleen extending 5½ inches below costal arch Many L D bodies found on spleen puncture Body weight—3 st 3 lbs Temperature ranged from 100°F to 102°F Bleeding from the gums and nose present Patient was treated with intravenous injection of urea stibamine, the dose being 5 c c of a 2 per cent solution (= 1 grm) During treatment patient developed cancrum oris Temperature began to come down after four injections and after six injections it remained permanently normal Altogether 13 injections were given after which the spleen could hardly be felt below costal arch Patient is still in hospital, 80 days after completion of treatment Increase of body weight—1 stone Patient's general condition—very satisfactory Duration of treatment—43 days

Blood count	Peripheral blood culture N N N medium	Spleen puncture	Spleen blood culture NNN med um	Remarks
R B C -2,800,000 W B C -1,000, Hb-42 % R B C -3,200,000, W B C -3,600, Hb-46 %	Positive Positive	Positive Positive	Positive Positive	Before treat ment 18 days after com mencement of treatment, 6 in jections having been given Total amount 6 grm

BI dc nt	P ph I II d ult NNN med m	Spl n pretr	Spl n bl d clt NNN m d m	Remarks
R B C -4 200 000 W B C -6 407 Hb-48	N B I	V & t	Ngt	32 dy ft omm n mnt f tr t m t 1011c to ha g b n g n T t l sm tl g m
R B C -46'00 10 W B C -70'00 Hb-48	NEI	Ngt	Ngt	d3 dy fi comm n m t ft t m t l3 jc to ha g b ng fit wh ch th yw topp d T t I m u t - 14 g m
R B C - 4 600 000 W B C - 9 000 Hi - 52°	Net	Ngt	Ngt	53 dy ft omplin of t tment patet till nhosp t l G I n dt — y t f ct y

Further Notes on Cases Previously Reported in the Indian Journal of Medical Research October 1922

No 1 Patient Monu left hospital 75 days after treat ment was stopped. At the time of discharge no L. D bodies were found on spleen puncture. Spleen could not be felt below the costal arch. Increase in weight—I stone. No fever since treatment was stopped. General condition—very satisfactory. R. B. C.—4.200.000. W. B. C.—8.600. Hb—54/ at the time of discharge.

No II Patient Tofu left hospital 51 days after treat ment was stopped. At the time of discharge no L D

bodies were found on spleen puncture. Spleen could not be felt below the costal arch. No fever since treatment was stopped. General condition very satisfactory. R. B. C.—4,400,000, W. B. C.—10,200, Hb—60% at the time of discharge.

No IV Patient, Anath Bandhu, left hospital 50 days after treatment was stopped. At the time of discharge no L. D. bodies were found on spleen puncture. Spleen could not be felt below the costal arch. No fever since treatment was stopped. General condition very satisfactory. Increase of weight—9 lbs. during 30 days after completion of treatment. R. B. C.—3,400,000, W. B. C.—8,200, Hb—58%

No VII Patient, Horoz, is still in hospital, ie, 10 months after treatment was stopped. Cultural reports are attached herewith. General condition very satisfactory. No fever since the treatment was stopped.

Blood count	Peripheral blood culture N N N medium	Spleen puncture	Spleen blood culture NNN medium	Remarks
R B C - 4,800,000, W B C - 7 200, Hb - 52%	Negative	Negative	Negative	130 days after treatment was stopped
R B C —5,100,000, W B C —8,600, Hb—60%	Negative	Negative	Negative	305 days after treatment was s t o p p e d Patient is still under observation in hospital, in very excellent condition of health

Remarks—Case No VII, Horoz, has been kept in hospital for a prolonged period to test the permanency of the efficiency of treatment. The remaining four cases reported in

my first paper and not reported here left hospital soon after my first paper was sent for publication

The method adopted for the culture of L donovani in my experiments is as follows (Major Knowles, 1 M S)—it is a modification of Row's method —

About a quarter of a c c of blood from a vein at the bend of the elbow is put into 20 c c of citrated salt solution (normal saline containing 1 5 per cent sodium citrate) the mixture is shaken gently and allowed to stand for some time. As soon as the corpuscles have settled to the bottom of the tube the supernatant fluid is poured off and the corpuscles are pipetted off with a capillary pipette and inoculated into the water of condensation at the bottom of N N N tubes which are then incubated at 22°C

In the case of splenic blood the syringe is filled with a few drops of the citrate solution mentioned above and then the spleen is punctured. The blood drawn is then mixed with the citrate saline inside the syringe transferred to the NNN medium and incubated in the same way as in the case of peripheral blood.

In an untreated case the parasites flagellate generally within the first week

NB—To ensure a successful result strict aseptic pre cautions are necessary as even slight bacterial contamination kills the LD parasites though they have been found to grow luxuriantly with fungi

I think Frankel is not justified in stating in his. Die Arzentimittel Synthese—that changes in the molecular structure of antimony compounds do not bring about an increase of their therapeutic properties—Urea stibamine is just as useful in the treatment of kala azar as atoxyl or soamin in diseases for which they have been recommended

REMARKS

A series of cases have been described which have been cured by the intravenous injection of urea stibamine

My grateful thanks are due to Major Knowles, I M S, Protozoologist, Calcutta School of Tropical Medicine, for helping me in making the cultural tests in his laboratory. I am also indebted to my staff, Mr. Saroda Churan Chowdhury, M Sc, Mr. Judhistir Dass, M Sc, Sub-Asst. Surgeons, Bibhuty Bhushan Maity and Sirish Chandra Banerjee, for their faithful services and hearty co-operation in helping me in carrying on my researches.

Through the kind courtesy of Major H E Shortt, I M S, Assam Scientific Research Committee, Shillong, I am appending here an extract from his notes on the use of urea stibamine—

EXTRACT FROM A NOTE ON THE USE OF UREA STIBAMINE

RY

MAJOR H E SHORTT, I M S Assam Scientific Research Committee, Shillong

Five cases altogether were treated with this preparation and, in our opinion, with most encouraging results. As the results obtained in some at least of these cases were very striking, the particulars of each case will be mentioned in some detail in the annexed table. The dosage employed by us was that recommended by Dr. Brahmachari, the solution for each injection being made up afresh. The initial dose was 0.1 gram dissolved in cold sterile distilled water. Each subsequent dose, given on alternate days, reached 0.25 gram which was not exceeded. Thus the fourth and all subsequent doses were of 0.25 gram. As the preparation is precipitated from alcohol, it is presumably sterile

and on solution in cold sterile distilled water needs only warming in a water bath. Administration was made by intravenous route. Results of spleen puncture were tested by microscopic and cultural methods. The details of the cases treated are given below in tabular form. The perusal of the table will at once show that the results obtained were so favourable as to encourage one to make a further extensive trial of this preparation. Its advantages over the antimony preparations usually employed as evidenced by experience of it in these five cases are

- (1) The short course occupying only two to three weeks necessary to a complete cure
- (2) The rapidity with which the symptoms of the disease disappear
- (3) The fact that no symptoms of intolerance were met with in any of the cases

The results here recorded are actually better than those claimed by Dr Brahmachari (1922) himself in the published account of some of his cases where he gave as many as 20 injections

TABLE Showing Results of Cases Treated with Urea Stibamine

	Remarks	Case on admission was very ill with marked ædema of legs and great weakness	Case on admission weak with cedema of feet Spleen puncture was not repeated before the eighth injection	Case on admission, emaciated, weak and very anæmic	Case on admission was extremely weak with severe bronchitis Parasites very numerous	Case had previously received full treatment with sodium antimonyl tartrate without benefit
	Result	cure	cure	cure	cure	
	Weight in pounds before and after treatment	1173-133	78-99	56-711	93-110	
	Total amount of urea strbamine administered	17 grammes	17 grammes	17 grammes	2 295 grammes	Still under treatment
(a atman = 6,	No of injections after which spleen puncture was negative	5	æ	4	6	4
G	Amount of urea stib-mine after which splech puncture was negative	95 gramme	17 grammes	7 gramme	l 795 grammes	7 gramme
2	Result of spleen puncture	+++++	+ + + +	+ + +	+ + +	+ +
	Duration of illness on admission	11 months	11} months	6 months	9½ months	12 months
	Age	72	17	15	40	38
	No No	43	48	95	57	58

CHEMOTHERAPY OF ANTIMONIAL COMPOUNDS IN KALA AZAR INFECTION

[Rece edf Publ ton D cembe 19 1922]

PART Y

AMINO ANTIMONYL TARTRATES

General Method of Preparation of the New Amino
Animonal Tartrales

- (1) In the first method molecular proportions of tartaric acid and the base were dissolved in boiling water. To this solution were subsequently added molecular proportions of antimony trioxide. The whole was heated till the oxide went into solution. The solution was then filtered and the filtrate solidified by gentle heating. The product was purified by repeated crystallization from water. In some cases the acid tartrate of the base was boiled gently with antimony trioxide.
- (2) In the second method molecular proportions of tartar emetic or sodium antimonyl tartrate and the hydro chloride of the base were refluxed in a medium of alcohol and water (6 to 2) for an hour. The solution was filtered and the filtrate was then gently heated till a solid mass was obtained. The salt was purified from sodium or potassium chloride by repeated crystallization from water and alcohol. The following reaction takes place.

C4H4O6 SbOk+R HCI=C4H5O6SbO R+KCI

(3) In the third method, molecular proportions of silver antimonyl tartrate and the hydrochloride of the base were either refluxed in a medium of water for an hour or kept at the ordinary temperature for a day with frequent and vigorous agitation. When the reaction was complete the solution was filtered and the filtrate concentrated, either in the water bath or in vacuo, when a solid or semi-solid mass resulted. In the first case, the mass was purified by recrystallization from water, while in the second case the semi-solid mass became solid on treatment with absolute alcohol. This solid mass was purified by dissolving in water and subsequent precipitation with alcohol or with alcohol and acetone. The following reaction takes place—

 $C_4H_4O_6$ OSbAg+R $HCI=C_4H_5O_6$ OSb R+AgCI

Acriflavine antimonyl tartrate was prepared in a special way to be described below —

(1) Preparation of p-phenetidinyl-acetamido-antimonyltartiate (= phenocoll antimonyl tartiate)

15 grms of tartaric acid were dissolved in water and to the solution were added phenocoll base 2 grms and the whole was heated till the base went into solution completely. Antimony trioxide 2 grms were gradually added to it and the whole was boiled for 15 minutes. It was then filtered, the filtrate being concentrated and the oil separated. This oil on cooling and after agitation solidified into a crystalline mass. This was then purified by double recrystallization from water.

Yield—almost theoretical

Calculated for $C_{14}H_{19}O_{9}N_{2}Sb$, Sb=25.06% Found Sb=25.08%

This compound has been prepared with the idea of using it intramuscularly

(2) Preparation of p carbethoxy aniline aniimanyl tartrate(=anæsthesin aniimonyl tartrate)

l 5 grms of tartaric acid were dissolved in water and to the solution were added l 6 grms af the ethyl ester of p amino benzoic acid and excess of water the mixture was then boiled until the whole mass went into solution. To this solution was added gradually antimony triaxide in excess (nearly l 6 grms) and the whole was boiled far half an hour lt was filtered and the filtrate on cancentration and cooling produced a crystalline mass. It was dissolved in water under reflux and allawed to cool when well defined crystals were obtained. (Yield=2 grams.)

Calculated for C13H16O9 Sb Sb=26669 Found Sb=266/

Praperties—It is not very freely aluble in water but its salubility is markedly increased in water cantaining a trace of tartaric acid

This campaind was prepared with the idea of using it internally as anæsthesin has a marked anæsthetic effect an the stamach

(3) Preparation of navacame antimanyl lartrales

Two antimonyl tartrates have been abtained fram navacaine in the first one molecule of antimanyl tartaric acid combined with novocaine base while in the second twa molecules of antimonyl tartaric acid combined with the base

These campounds have been prepared with the idea of using them inframuscularly

(a) Preparation of novocaine mano antimonyl tartrate

0 546 grm of novocaine hydrochloride was dissolved in 20 c c of water in a 50 c c flask and to this was added 0 82 grm of silver antimonyl tartrate The flask was well corked protected from sunlight by non-actinic paper and frequently vigorously shaken. It was kept in this condition for six days after which it was filtered. The filtrate was tested and found free from chlorine and silver. The filtrate was allowed to concentrate in a vacuum desiccator. On concentration the solution set to a gel which on treatment with alcohol yielded a white solid. This was quickly collected, washed with ether, and again dried in a vacuum desiccator.

Calculated for $C_{17}H_{25}O_{9}N_{2}$ Sb. $(C_{2}H_{5})_{2}$ CH₅O₆ (SbO), $S_{\nu}=23~03\%$ Found Sb=23% Yield= 3 grm

It is highly hygroscopic and is easily soluble in water

(b) Preparation of novocaine-di-antimonyl tartiate

15 grams of tartaric acid were dissolved in 30 c c of water by heating. To the solution were added 12 grms of novocaine base and then gradually 16 grms of antimony trioxide while the whole was kept boiling for half an hour after addition of antimony trioxide. The unattacked antimony trioxide was separated by filtration, the filtrate concentrated and absolute alcohol added to the solution, when a white crystalline highly hygroscopic substance was obtained. This was next treated with alcohol and acetone and subsequently kept in a vacuum desiccator over sulphuric acid.

Calculated for $C_{21}H_{30}O_{16}N_2Sb_2$, Sb=29.7% Found Sb=29.2%

Novocaine-di-antimonyl tartrate is highly hygroscopic and undergoes decomposition when kept in a wet condition in the air, yielding a yellow substance — It is very soluble in water

(4) Preparation of apothesine antimonyl tartrate

EXPERIMENTAL

1 2 grms of apothesine hydrochloride are dissolved in water and to the solution are added 1 64 grms of silver anti-

monyl tartrate After 48 hours during which it is frequently and vigorously shaken the mixture is filtered when the fil trate was found to be free from chlorine and silver. It is then placed in a vacuum desiccator and on concentration a jelly like mass was obtained which solidified on treatment with absolute alcohol. It is collected and washed with absolute alcohol and acetone and dried in a vacuum desiccator (Yield = 0.5 gram.)

Calculated for C 0H 8O9NSb Sb=21 97 / Found Sb=21 96 /

The salt is highly hygroscopic and is easily soluble in water

The compound was prepared with the idea of using it intramuscularly

The toxicity of two of the above antimonyl tartrates is given below —

Lethal effects obtained from the administration of a one per cent solution of phenocoll antimonyl tartrate into guinea pigs by intramuscular injection

D gmpklo	Nofnmlud	N mb d d
0.08	3	3
C 075	3	2
0 07	3	7
0 065	3	2
0 055	1	0

TABLE I

Toxicity of the antimony content = $\frac{k}{0.08 \times 25.06}$ or $\frac{k}{2.0048}$

Lethal effects obtained from the administration of a one per cent solution of anæsthesin-antimonyl tartrate into guineapigs by intramuscular injection

T	ABLE	Π

Dose in grams per kilo	No of animals used	Number died		
0 09	3	3		
0 085	2	1		
0 08	1	0		

Toxicity of the antimony content =
$$\frac{K}{09 \times 26.6}$$
 or $\frac{K}{2.394}$

(5) Preparation of methyl-amino-hydroxybenzoate antimonyl taitiate (= oithoform antimonyl tartrate)

EXPERIMENTAL

15 grms of tartaric acid dissolved in 50 c c of water are heated with 17 grms of 3-amino-4-hydroxy-methyl benzoic ester till the latter goes into solution. This solution is filtered and the filtrate is boiled for nearly two hours with 16 grms of antimony trioxide. The filtrate on being concentrated and the side of the basin being scratched, a slightly yellow tinged substance separates. It is filtered and purified by crystallization twice from water. (Yield=2 grams.)

This compound has been prepared with the idea of introducing antimony through the abraded skin or as an inunction

Calculated for $C_{12}H_{14}O_{10}$ NSb, Sb = 26.54% Found Sb = 26.25%

(6) Preparation of 3 6 diamino 10 methyl acridinium anlimonyl tartrate (= acriflavine anlimonyl tartrate)

0 65 grm of 3 6 diamino 10 methyl acridinium chlo ride is dissolved in 20 cc of pyridine and the solution filtered. To the filtrate is added a solution of 1 grm of silver antimonyl tartrate in 20 cc of pyridine. Reaction takes place at once and a light yellow precipitate is obtained. The whole mixture is then refluxed for at least four hours and the precipitate that is formed is collected by filtration. It is purfied by washing thrice in hot pyridine, then several times with hot alcohol and subsequently with distilled water.

Calculated for $C_{18}H_{18}O$ SbN, Sb=23 62% Found Sb=23 92% The compound is sparingly soluble in water and is lighter in colour and less fluorescent in solution than acriflavine

This compound was prepared with the idea of combining the therapeutic properties of antimony and acriflavine

REMARKS

The late Sir Patrick Manson once wrote to me as follows. Go on in your efforts to get an antimony compound that can be used as an intramuscular injection or better still as a drug that can be administered by the mouth. It now requires to be seen how far the inference of the proper ties of the above antimonyl tartrates based on theoretical considerations is borne out in practice and how far they exert leishmanocidal properties. I shall enter into the subject in a subsequent series.

CHEMOTHERAPY OF ANTIMONIAL COMPOUNDS IN KALA-AZAR INFECTION

[Received for Publication, March 13, 1923]

PART VI

CUMULATIVE AND TOLERANCE EXPERIMENTS WITH TARTAR EMETIC

THE following series of experiments have been made to determine whether repeated injections of tartar emetic into guinea-pigs in sub-lethal doses lead to tolerance towards the drug or its cumulative action

	Injections		Dose in	Body weight	Injections		Dose in	Body weight
Serial No	No	Date	per kilo of body weight		No	Date	per kilo of body weight	in grms at time of injuc tion
	1 _{st}	25 8 22	005	340	2nd	28 8 22	0075	355
	3rd	31 8 22	01	360	4th	4 9 22	015	355
771 {	5th	7 9 22	02	360	6th	12 9 22	025	380
(/th	18 9 22	03	385	8th	22 9 22	035	360

Remarks-Animal died on 22nd September, 1922 with post mortem signs of antimony poisoning Total quantity injected- 1475 grm per kilo

						 		· · · · · · · · · · · · · · · · · · ·
(20	İst	29 3-22	018	355	2nd	30 3 22	018	340
629	3rd	31 3 22	018	350	4th	1 4 22	018	355

Remorks—Death on 1st April, 1922, showing post mortem signs of antimony poisoning Total amount injected— 072 grm per kilo

564				ſ	2nd	7 4 22	018	250
J04 {	3rd	8 4 22	018	255				

Remarks—Death on 10 April, 1922, showing post mortem signs of antimony poisoning
Total quantity injected— 054 grm per kilo

ŧ

s	IN	Injections		Ds Bdy w ght	ÎNJE	CTIONS	Do g m	Eody w ght	
		N	Dt	p k lo f body w ght	ngm ttm of nj t n	N	Dt	p kl fb dy w ght	ngms ttme f j ton
	ſ	1 t	26 7 21	0°25	398	2 d	8821	0025	392
	ı	3 d	11 8 21	003	412	4th	19821	004	415
143	ł	5th	24 8 21	893	437	6th	30 8 21	10	435
		7th	5 9 22	01	41	8th	15 9 21	012	435
	į	}	16921		415	ļ	21921		425

Rm k -D th n 22 dS pt mb 1922 h wag p tm tm ga of at mo y p s g T tal q t ty njet d-052 pe k lo

	(1:	12 11 21	NO5	195	2nd	15 11 21	005	196
	3 d	19 11 21	007	210	4 h	21 11 21	007	200
	5th	23 11 21	008	210	61h	30 11 21	800	210
	7th	4 12 21	91	210	8th	7 12 21	01	210
	9th	11 12 21	015	230	10th	13 12 21	015	240
254	11th	16 12 21	02	265	121h	19 12 21	02	260
4)7	ii .	20 12 21		260	13th	22 12 21	025	250
	14th	25 12 21	025	275	15th	28 12 21	03	262
	16th	31 12 21	04	280	17th	3 1 22	045	292
		5 1 22		285	18th	8 [22	05	295
	19th	t5 1 22	06	320	1	20 1 22		322
	20th	21 1 22	07	330	ĺ	1		1

Rm k—Slightlyr ti detof dir mwhidll nd ymuch t feed dir whidll nd ymuch t feed dir whidll nd ymuch t feed no n Zand D c li li n n zah D mb 1921 A m lbecm dull 21; nury 1922 in th fin a add d taght howing p tm tm g of tm nypo n g T tild fit m t jet dip kio f bdy w ght—475 gmm

	Injections		Dose in	Body weight	Inje	CTIONS	Dose in grms	Body weight
Serial No	No	Date	per kilo of body weight	at time of inject tion	No	Date	of body weight	at t me of injec tion
				\ <u>-</u>	}			
ĺ	lst	5 9 22	0075	250	2nd	8 9 22	01	265
70:	3rd	13 9 22	015	280	4th	18 9-22	02	270
781	5th	22 9 22	025	275	6th	6 10 22	03	265
ł	7th	10 10 22	035	280				

Remarks—Animal became dull 6 hours after 5th injection. It began to lose weight
Animal died 10 hours after the last injection with post-mortem signs of
antimony poisoning. Total quantity injected—1425 gim. per kilo of
body weight.

{	lst	27 3 22	01	300	2nd	31 3 22	015	290
					4th	7-4 22	02	315
540	5th	12 4 22	025	315	6th	19 4 22	03	330
{	7th	26 4 22	04	330			7	

Remarks—Somewhat restless after injections on 7th and 12th April, 1922. The animal began to lose weight considerably after injection on 19th April, 1922, but regained weight after 6 days. Restless after the injection on 26th April, 1922. The animal died 12 hrurs after injection showing post moitem signs of antimony poisoning. Total dose of tartar emetic injected per kilo of body weight— 158 grm.

253	ĺ]		l i	15 11 21 21 11 21	217
)	25 11 21	1		2. 1, 2,	

Remarks—Died on 25th November, 1921, showing post moitem signs of antimony poisoning Total dose of tartar emetic injected per kilo of body weight— 032 grm

543		27 3 22						310
243	(3rd	3 4 22	018	330	4th	7-4 22	02	275

Remarks—Restless after injection on 7th April, 1922, and died the same night showing post mortem signs of antimony poisoning Total dose of tartar emetic injected per kilo of body weight— 063 grm

	1 дестючя		p t kilo	Body w ight	INJECTIONS		D se in g m pe kilo	Body weight	
Se i I No	No	Dι	of body w iht	at tim of Injec tien	٨	D te	of body weight	fijec fije	
			1		! -				
!	111	17722	05	370		13 7 22		360	
i	2nd	16 7 22	005	370		177_2		365	
ļ	3 4	18 7 22	01	365	4:h	20 7 22	01	365	
ļ		22 7 22		367	541	24 7 22	10	3.4	
747	6th	27 7 22	015	3,2	7d	31 , 22	10	365	
,	8:h	3 8 22	C15	360	916	6822	75	350	
	10th	9822	035	355	160 '	12 8 72	04	156	
	12 h	15 8 22	1 (45	369	134	20 8 22	05	370	
	14th	75 8 22	055	357	15d	25 8 22	755	312	

Rm ku-Somewhatril o 17th Jly 1922. The milimid dill for two dy firliecti on 20th Agri 1922. The imild don 28th Agri 1922. Shou firliection Totld Itrm tic inject dorikil of body wight-405 gm.

	1 1:	15 7 22	005	325	2 d	1872	005	325
48	34	20 7 22	0975	325	4:1	24 7 22	01	42
	!! !	29 / 22		310	બા	31722	01	315
	6th	6822	02	32D		7 8 22		312
40	7el	9 8 22	025	335		10 8 22		320
	8th	12 8 22	03	340	9tl	16 B 22	035	340
	il l	18 8 22		307	10:h	23 8 22	04	320
	11th	25 8 22	0045	325	17th	28 8 22	05	25

R m k -- 12

	Injections		Dose 111	Body weight	Injections		Dose in grms	Body weight
Serial No	No	Date	grms per kilo of body weight	in grms at time of injec tion	No	Date	per kilo of body weight	in grms st time of injec tion
(lst	25 8 22	005	402	2nd	28 8 22	0075	390
	3rd	31 8 22	01	410	4th	5 9 22	015	390
772		7 9 22		375	5th	8 9 22	02	335
	6th	13 9 22	025	405	7th	18 9 22	03	405

Remarks — Died on 19th September, 1922, showing post mortem signs of antimony poisoning Total dose of tartar emetic injected per kilo of body weight—
1125 grm

	İst	5 9 22	0075	415	2nd	8 9 22	01	430
	3rd	11 9 22	015	425	4th	18 9 22	02	428
	5th	22 9 22	025	445		23 9 22		425
	,	6 10 22		430	6t h	10 10 22	035	450
782		11 10 22		435		14 10 22		430
į	7th	15 10 22	04	440		16 10 22		430
]		18 10 22		430		19 10 22		420
Ì	8th	20 10 22	045	390	9th	30 10 22	05	390
l	10th	12 11 22	055	365				

Remarks — The animal became dull 12 hours after injection on 22nd September, 1922. The animal became dull after injection on 15th October, 1922, and gradually lost weight since. Death took place 8 hours after last injection. Total dose per kilo of body weight— 3025 grm.

Conclusion

I have observed that repeated injections of tartar emetic in sub-lethal doses did not give rise to any tolerance towards the drug except very rarely. Generally the results pointed to a cumulative action of the drug, or at least made the animal susceptible to the next higher dose.

CHEMOTHERAPY OF ANTIMONIAL COMPOUNDS IN KALA AZAR INFECTION

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PART VII

DETERMINATION OF SMALL QUANTITIES OF ANTIMONY IN PRESENCE OF ORGANIC MATTER

The method followed by us in the quantitative determination of antimony in stools and urine depends upon the extraction of the antimony by boiling with eopper and hydro chloric acid and the subsequent solution of the antimony and its conversion into sulphide which is estimated colori metrically. We at first tried the method described by Schidrowitz and Goldsbrough (Analyst 1911 36 101) and its modification by B am and Freak (Analyst June 1919). Both these methods gave unsatisfactory results in our hands and the difficulties that we had to encounter were due to the presence of iron and the use of alkaline permanganate solution.

(1) In the process of extraction of antimony from urine minute traces of iron are frequently deposited over the eopper from the urine and go into solution in the subsequent treat ment with alkaline permanganate solution. The iron seems to be converted into a soluble ferrate from which it cannot be precipitated by the alkali in the alkaline permanganate solution. Beam and Freak's procedure does no remove this iron which passes into the final test solution giving rise, in the presence of antimony to a reddish brown colour with

hydrogen sulphide which cannot be made to match with the yellow colour of pure antimonious sulphide solution

- (2) It seems that it is extremely difficult to prevent some manganese from passing into solution and its presence interferes considerably with the colorimetric test
- (3) As pointed out by Beam and Freak, the antimony precipitated on the copper becomes difficult to dissolve if it is not immediately treated with alkaline permanganate, and prolonged treatment with alkaline permanganate leads to the solution of copper

We have therefore modified Beam and Freak's procedure to a considerable extent and our method is described as follows —

- (1) The whole of the twenty-four hours' urine is concentrated in a porcelain dish to about 50 c c by heating. The concentrated urine is then mixed with 10 c c of chemically pure concentrated hydrochloric acid and strips of pure copper foil of suitable size are introduced into the solution, which is boiled for some hours till all the antimony is deposited over the copper. To ensure complete precipitation of the antimony, a fresh strip of copper foil is introduced into the solution which is again boiled, after addition of a fresh quantity of chemically pure hydrochloric acid and distilled water, till no more black deposit forms on the copper. The strips of copper are then removed with glass forceps and, after washing in distilled water, are treated with an alkaline solution of persulphate of potassium
- (II) After the antimony has completely gone into solution, the latter is boiled with a slight excess of alkali to precipitate any traces of copper that may have gone into solution
- (III) The solution is filtered and sulphur dioxide passed into it for three to five minutes. The solution is boiled with hydrochloric acid which converts the iron into a chloride

- * (IV) After expelling the hydrochloric acid and the sulphur dioxide the solution is boiled with potassium hydrate to precipitate the whole of the iron and is then filtered
- (V) The filtrate is acidified with hydrochloric acid and then a sufficient quantity of distilled water is added to make it up to 100 c.c.
- (VI) To the solution 5 cc of 10 per cent gum olution is added and hydrogen sulphide passed through 50 cc of the solution until the colour is fully developed. Comparison is then made with a standard solution of pure tarter emetic treated in the same manner in Nessler's tubes and the amount of antimony thereby estimated.

To test the accuracy of the procedure the following estimations were made with chemically pure tartar emetic (sample supplied by Messrs Martindale & Co and certified by them to be one hundred per cent pure)

	Solut on 1 ken	Atmonyf nd
(1)	0009 grm of antimony	00087
(2)	00144	00141
(3)	CO18	00177
(4)	0027	00267

In estimating antimony in the faces the stools are first mixed with water and boiled gently with hydrochloric acid for some hours and filtered. The faces are then extracted with hot water several times till no more antimony can be detected in the filtrate. The whole of the filtrate is now treated in the same way as above to extract the total amount of antimony.

REMARKS

- (1) The use of alkaline persulphate avoids the introduction of manganese into the final test solution, the presence of which interferes with the colorimetric test to a considerable extent.
- (2) Alkaline persulphate dissolves the deposit of antimony much more quickly than alkaline permanganate
- (3) It is essential to get rid of all traces of iron that may be precipitated from the urine on the copper foil and subsequently pass into solution
- (4) The method advocated here gives a fair degree of accuracy in estimating minute traces of antimony

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CHEMOTHERAPY OF ANTIMONIAL COMPOUNDS IN KALA AZAR INFECTION

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PART VIII

QUANTITATIVE STUDIES IN EXCRETION OF ANTIMONY (TARTAR EMETIC AND URFA STIBAMINE)

The excretion of antimony after administration of an antimonial to man or animals in the form of an inorganic or organic antimonial compound has received very little attention though considerable attention has been given to the excretion of arsenic since the introduction of arseno benzol compounds into therapeutics. Yet the former is just as important as the latter since the discovery of the very important part played by antimony in therapeutics.

The antimonials that have been used for the deter mination of the rate of excretion of antimony consist of antimonyl tartrates and urea stibamine. Of the antimonyl tartrates the rate of excretion of antimony after administration of tartar emetic will form the subject matter of the present paper. The antimonials studied here may be grouped together under the following heads. (1) Antimonials containing pentavalent antimony e.g. urea stibamine. (2) antimonials containing trivalent antimony—the antimonyl tartrates. We did not study the rate of excretion of antimony after administration of stibacetin (Stibenyl), as the drug has not given satisfactory results in the treatment of kala azar in the hands of many observers and just as atoxyl or soamin has replaced are acetin in the treatment of diseases where one

intends to use pentavalent arsenic, so the salts of p-amino-phenyl-stibinic acid will perhaps replace stibacetin. (Anti-monials of the stibinobenzene type have not yet come into use in the treatment of human diseases, though they have been used with indefinite results in the case of certain diseases of animals (vide Frankel's Arzentimittel Synthese)

By studying the excretion of antimony we have come to the conclusion that antimonials of the type of urea stibamine are converted within the body into trivalent oxide of antimony, similar to what happens in the case of organic pentavalent arsenicals, before they are capable of exhibiting organotropic and parasitotropic properties. This may be illustrated graphically in the following way—

$$\begin{array}{ccc} OH & \\ R & Sb & = O \\ OR^{t} & \end{array} \longrightarrow \begin{array}{c} R & Sb = O \end{array}$$

It is possible that -Sb=O is more reactive and more powerful in its parasiticidal properties when an organic antimonial is changed within the body into one containing it than when it exists in the antimonyl tartrates

We may assume that the parasiticidal properties of antimony compounds depend upon the radicle -Sb=O, similar to what exists in the case of arsenicals, in which the parasiticidal properties depend upon -As=O'

In the present paper, our experimental work on the excretion of antimony is limited to observations in man Though in some respects experiments on animals may be more convenient, yet the experiments conducted on man have the obvious advantage of giving an idea of the rate of excretion in man directly without the possible objection of having to deduce the same indirectly from observations on animals

Our procedure was as follows The individual, generally an adult, was asked to empty his bladder immediately before

injection and was injected intravenously generally between 9 A M and 10 A M daily. He was made to pass all his urine during 24 hours in perfectly clean bottles on the days of observation. The stools were similarly collected in perfectly clean vessels.

The same methods were adopted in the detection of antimony in the urine and the stools as were followed in its detection in the viscera (see Part III). The method adopted for the quantitative determination of antimony excreted has been described in the previous paper (see Part VII). As stated therein the methods that have been described by previous workers for the quantitative esti mation of antimony are misleading and we had to modify them considerably in order to obtain accurate results. It may be stated here that the estimation of antimony in the excrete tasks the patience of the experimenter to the utmost requiring the most careful and delicate handling of all the processes. All the reagents and water u ed in the experiments must be previously tested to ensure that they are free from arsenic or antimony.

TARTAR EMETIC Period of Presence of Antimony in Urine

(a) After Single Injection

			P	dd gwhh ntmnyw pr tnthun
1) 0	d of 25	f 2 p nt	Itng not vnuly	1 560 h u
(2)	D tt	Dι	D tto	774
(3)	D tt	D tto	D tt	ī 108
(4)	D tt	Dtt	D sto	1 060
		А σь	ar 4_46.4	

(b) After Repeated Injections

	eriod during which antimony vas pre sent in the urine
(1) 42cc of 2 per cent solution given in 15 injections (81 cm)	1 152 hours after the last injection
(2, 10 c c of 2 per cent rolution in 5 injections (2 km)	1 102 hours after the last injection

Average period -19 days

Daily Excietion of Antimony in Urine

(I)	(2)			(3)		i	14		
Injection of 2.5 c c of 2 per cent solution (= 018 Sh)	Injection of 10 of 2 per cent rolution (= 072)		of	on of 2: 2 per cer n (= 0 18	1t	of 2	tion of percent (036	rolu	
1 00108 Sb	96، د ۱ ا	Slo	r 1	01206	51,	1	002115	s _b	
2 00108 Sh	2 003120	Sh	2	00108	Sb	2	00218	Sb	
3 00099 Sb	3 002880	S1,	3	00000	Sb	3	001(71	Sh	
4 00054 Sb	1 002562	Sb	3	00003	Sb	4	02117	Sb	
3 000321 Sb	5 001800	Sb	5	00045	Sb	5	00108	Sb	1
6 000324 Sb	6 001480	Sb	6	09035	Sb	6	00108	Sb	
	7 001170	Sb				7	ტები	Sb	
	8 000972	Sb	t						
	9 000729	Sb							
	10 000648	Sb							
	11 000657	Sb							
	12 000648	Sb							

1st Observation -

Quantity of Sb passed in 24 hours $=\frac{1}{16.6}$ of the quantity injected

Quantity of Sb passed in 6 days = 004338 grm or $\frac{1}{41}$ oft he quantity injected

2nd Observation -

Quantity of antimony passed in 24 hours $=\frac{1}{18.5}$ of the quantity injected

Quantity of antimony passed in 6 days = 016138 grm or $\frac{1}{4.5}$ of the quantity injected

Quantity of antimony passed in 12 days = 020962 grm or $\frac{1}{34}$ of the total quantity injected

3rd Observation -

Quantity of antimony passed in 24 hours $=\frac{1}{14}$ of the quantity injected

Quantity of antimony passed in 6 days = $\frac{1}{38}$ of the quantity injected

4th Observation -

Quantity of antimony psssed in 24 hours = $\frac{1}{16.2}$ of the quantity injected

Quantity of antimony passed in 6 days = $\frac{1}{3.8}$ of the quantity injected

UREA STIBAMINE

Period of Presence of Antimony in the Urine

(a) After Single Injection

					dd ngwhh ntm yw p t thu
(I) O d	F 2 5	f2p c	t lut ng	t ıly	744 h r
(2) D	tt	D tt	D tto	Dtt	768
(3) D	tto	Dtto	D tt	Dtt	t 224
(4) D	tŧ	Dtt	Dn	Dtt	1 160
		A	g -40 dy		
15-76	57B				

(b) After Repeated Injections

- (1) 135 cc of 2 per cent rolution fixen in 14 injections (2.7 great) 1,537 hours
- (2) 100 cc of 2 per cent volution fisch in 15 injectio in 12 0 first 769 ...
- (3) 975 cc of 2 per cent solution given in 15 injections (10) prime 643

Average - 41 days

Daily Exerction of Antimony in Urinc

1

2

Injection of 5 c c of a 2 per cent colution	Injection of Sec. of 2 per cent will tion
= 037 gim Sbi	/ 037 sin
1 0110 grm Sb	1 01108 cm Sb
2 00189 grm Sb	2 (4)18 prm Sb
3 001 grm Sb	3 M126 cm Sb
4 0009 grm Sb	1)
5 0004 grm Sb	5 00141 grm St
6 00054 grm Sb	6 00072 prin 91,
	7 00036 gim Sb
	8 0593 gim Sb

1st Observation -

Quantity of Sb passed in 24 hours = 0149 Sb or $\frac{1}{2.5}$ of total quantity injected

Quantity of Sb passed in 6 days = 01963 grm Sb or $\frac{1}{19}$ of the total quantity injected

2nd Observation —

Quantity of Sb passed in 24 hours = $\frac{1}{3}\frac{1}{34}$ of the quantity injected

Quantity of Sb passed in 6 days = 0163 gim Sb or $\frac{1}{2.24}$ of the total quantity injected

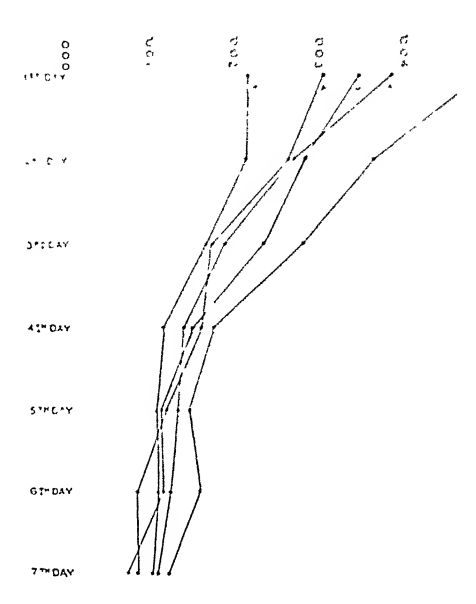
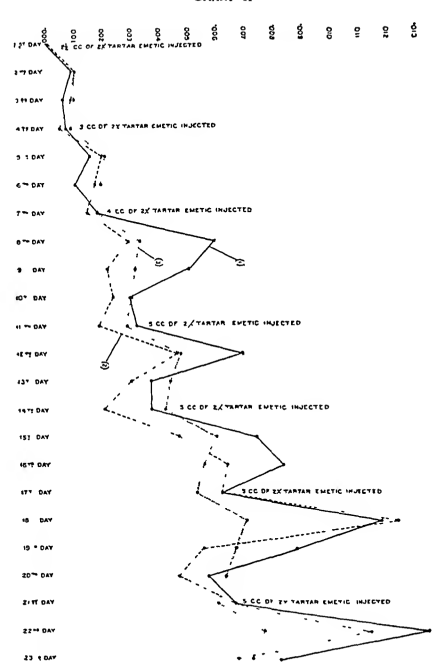


Chart showing the curve of excretion of antimonyl tartrates after intravenous injection in terms of antimony

[Reprinted from the Indian Journal of Medical Research, Vol XI, No 3, January, 1924]

CHART II



- l and 2 Curves of excretion of tartar emetic in terms of antimony after repeated injections of tartar emetic (2 cases)
- 3 A theoretical curve showing the excretion of tartar emetic in terms of antimony after repeated injections if the rate of excretion always followed the law that the amount excreted was proportional to the amount present in the system

We have ob erved that there is no relationship between the quantity of urine passed during a definite period and the excretion of antimony during the same period as will be seen by the following figures

Curve of Excretion of Antimony in the Urine

In the first observation after injection with tartar emetic the total quantity of urine passed in 24 hours was 590 c c and in 6 days it was 3 097 c c. Amount of Sb pas ed in 24 hours = $\frac{1}{16.6}$ and in 6 days $\frac{1}{4.1}$ of the total quantity injected

In the second observation after injection with tartar emetic the total quantity of urine passed in 24 hours was 1 625 c c and in 6 days it was 12 625 c c. Amount of Sb passed in 24 hours was \frac{1}{185} and in 6 days was \frac{1}{45} of the total quantity injected. The quantity of urine passed by the first case during the first 24 hours. Similarly the quantity passed during the 6 days by the second case was four times that passed by the first case. The same phenomena were also observed in the case of urea stibamine.

Thus in the first observation the amount of antimony passed in 24 hours was $\frac{1}{2.5}$ and in 6 days $\frac{1}{1.9}$ of the antimony injected. The amount of urine passed in 24 hours was 1 650 c.c. and the amount passed in 6 days was 8 225 c.c.

In the second observation the amount of antimony excreted in 24 hours was $\frac{1}{34}$ and in 6 days it was $\frac{1}{227}$ of the total quantity injected. The total quantity of urine passed

during 24 hours was 440 c c., being nearly one-fourth the quantity passed in the first case During 6 days, the amount passed was 2,665 c c, being less than one-third the amount passed by the first case during the same period

It will also be observed that in the case of tartar emetic the curve of excretion is one slowly converging to the base line, while in the case of urea stibamine the curve is abrupt during the first 24 hours and then follows the same course as in the case of tartar emetic.

It is evident that the excretion of antimony, after injection of urea stibamine, during the first 24 hours is quick and follows a similar course to what is observed in the case of organic pentavalent arsenicals, and then follows a curve like that in the case of tartar emetic which contains trivalent antimony. From this one may conclude that in the excretion of pentavalent antimonials, antimony is excreted as a pentavalent antimonial compound during the first 24 hours and then undergoes a reduction of the nature described before and in this process of excretion gives rise to a compound containing—Sb=O in a reactive stage

Excretion of Antimony in the Fæces and Vomit

The excretion of antimony in the fæces is not so regular as that in the urine

TARTAR EMETIC

The excretion after intravenous injection of 10 c c of a 2 per cent solution of tartar emetic was as follows —

(1) 1st 24 hours—no stool (2) 2nd 24 hours— 001008 grm Sb (3) 3rd 24 hours—no stool (4) 4th 24 hours— 000936 grm Sb (5) 5th 24 hours—no stool (6) 6th 24

hours— 00072 grm Sb (7) 7th 24 hours—stool too small to be quantitatively determined

Total quantity = 002664 grm Sb or $\frac{1}{27}$ of the total quantity injected

After intravenous injection of 2 5 c c of a 2 per cent solution of tartar emetic the amount of antimony present in the stools from the very beginning was too small to be quantitatively determined

UREA STIBAMINE

The excretion after injection of 5 c c of 2 per cent solution was as follows

(1) 1st 24 hours—no stool (2) 2nd 24 hours—00072 grm Sb (3) 3rd 24 hours—00054 grm Sb (4) 4th 24 hours—no stool (5) 5th 24 hours—00018 grm Sb (6) 6th 24 hours—too small 10 be quantitatively determined

Total quantity = 00144 grm Sb or $\frac{1}{25}$ of the total quantity injected

In another observation the total quantity of antimony passed in the fæces after injection of 5 c c of 2 per cent solution was 00063 grm or less than $\frac{1}{60}$ of the total quantity injected

In the vomit immediately after the intravenous injection of tartar emetic no antimony could be detected confirming the observations of Weiss and Haicher

REMARKS

1 The amount of antimony excreted in the urine during the first 24 hours after intravenous injection of tartar emetic is about 6 per cent of the amount injected

- 2. The amount of antimony excreted in the urine during the first 24 hours after intravenous injection of urea stibamine is 30 to 40 per cent. of the amount injected
- 3 The amount of antimony passed by the kidneys during the first 24 hours is fairly proportional to the amount injected.
- 4 There is no relationship between the quantity of urine passed during a certain period and the excretion of antimony during the same period. The effect of strong diuretics has not yet been studied
- 5 The excretion of antimony after intravenous injection of tartar emetic follows a curve slowly converging to the base line
- 6 The excretion of antimony after intravenous injection of a pentavalent organic antimonial follows a curve, the first portion of which, representing the excretion during the first 24 hours, is abrupt and the second portion follows a course similar to that found in the case of tartar emetic.

It is probable that a pentavalent organic antimonial is converted in the body into a trivalent antimonial and that as long as it exists in the body in the pentavalent form its rate of excretion is much quicker than when it is converted into the trivalent form. During the latter stage the curve of excretion is similar to that of tartar emetic in which antimony exists in the trivalent form.

- 7 After a single intravenous injection of an antimonial compound, the antimony may be present in the urine even for two months
- 8 Since a great portion of antimony present in an aromatic pentavalent antimonial (urea stibamine) is quickly eliminated, the chances of toxic action of the compound are much less than that of an antimonyl tartrate
- 9 In the process of conversion of an aromatic pentavalent antimonial in the body into a compound containing trivalent antimony, a reactive—Sb=O is formed, which is

probably responsible for the remarkably beneficial results observed in the treatment of leishmaniasis by the use of urea stibamine

- 10 The excretion of antimony by the fæces after intravenous injection of tartar emetic or a pentavalent auti monial is irregular and the quantity excreted is considerably less than in the urine
- 11 The low toxicity of urea stibamine the fact that no intolerance towards the drug is likely to take place on account of the quick elimination of a large proportion of it the fact that it is perhaps converted in the body into a trivalent antimonial containing a reactive—Sb=O radicle and the fact that its therapeutic value is very great (vide Chemotherapy of Antimonial Compounds in Kala azar Infection Parts I and IV in Indian Journal of Medical Research October 1922 and 1923 and also Major Shortt's paper in the Indian Medical Gazette July 1923 and in Indian Journal of Medical Research October 1923) make urea stibamine the best of all the antimomal compounds that have so far been discovered for the treatment of leishmaniasis and other diseases in which antimony is indicated

The relation between excretion of antimony and renal functional activity clinical character of the urine and physical and chemical characters of antimonial compounds used will be discussed in another series

CHEMOTHERAPY OF ANTIMONIAL COMPOUNDS IN KALA-AZAR INFECTION

[Received for Publication, September 17, 1923]

PART IX

TREATMENT OF CASES OF KALA-AZAR RESISTANT
TO ANTIMONYL TARTRATES WITH UREA
STIBAMINE—THE THERAPEUTIC VALUE
OF STIBAMINE IN KALA-AZAR

(A)

The first series of kala-azar cases treated with urea stibamine appeared in the Indian Journal of Medical Research in October, 1922. The subsequent researches of Shortt have confirmed my observations and proved the remarkable leishmanicidal properties of the compound (Indian Medical Gazette, July, 1923). My further observations and those of Major Shortt on its use appeared in the Indian Journal of Medical Research, January, 1924. The following are Major Shortt's notes on his three recent cases cured by urea stibamine.

1st case—Total	urea	stibamine	given	9	gramme	ın	5
					injections		
2nd case—	Ι) _o		75	gramme	ın	4
					injections		
3rd case—	Ι)o		65	gramme	ın	4
	injections						

In each case the cure was established by subsequent observations in hospital and negative results obtained by culture of puncture material

Writing to me recently on the use of this compound in kala azar Capt C Martin IMS of the Lawrence Military School Sanawar Simla Hills stated as follows

I am writing to tell you that the urea stibamine you sent me has worked wonders with one of my ca es which was not doing at all well with tartar emetic

The compound is now having an extensive use in the wards of Lt Col Barnardo CIE CBE IMS Principal Medical College Hospitals Bengal

In the present paper is recorded the value of urea stibamine in a series of refractory cases of the disease

By refractory or resistant cases of kala azar I mean cases which have resisted treatment with two grammes or more of sodium or potassium antimonyl tartrate given intravenously in the routine form of treatment of the disease extending over a period of two and a half months to three months or more It is well known that a certain percentage of cases are not cured or sometimes not even benefited by this dose of these antimonyl tartrates Experience has also shown that all of these residual cases are not absolutely refractory as some of them may be cured by a further course of treatment with antimonyl tartrates Generally speaking however symptom of intolerance towards the drug appear in many cases after two grammes of the salt have been administered intravenously. Cases that have had this dose and show clinically little or no improvement and show the presence of leishmania in their tissues will be regarded as resistant or refractory for the purpose of this paper

I have adopted this definition as some time ago certain observations conducted in Shillong led to the conclusion that all cases of kala azar were curable with 2 grammes

of tartar emetic. Subsequent observations did not confirm this view, as 34 out of 50 cases in Shortt's series were not cured with this dose. Out of these 34, 25 were subsequently cured by further treatment with sodium antimonyl tartrate for more or less prolonged periods, leaving a residual number of 9 which either died or left hospital uncured in spite of further treatment with sodium antimonyl tartrate In Mackie's statistics, 13 out of 20 cases were not cured with this dose. Out of these 13. 4 were cured by further treatment with sodium antimonyl tartrate and Mackie considered that one was wholly refractory, the case being a boy of 10, showing living parasites after administration of 379 c c of one per cent solution of the salt, which would be equivalent to nearly 4 grammes of the antimonial I shall, however, presently show that a case that resisted this dose was subsequently cured with a course of treatment with urea stibamine In my own statistics in a series of 200 cases I have records of nearly 60 cases which resisted treatment with 2 grammes of sodium or potassium antimonyl tartrate or both combined, and 20 were not cured by 6 grammes of these salts Generally speaking, it may be stated that cases that are not cured by 2 grammes of sodium or potassium salt require very prolonged treatment with these salts and even then, some may not be cured or benefited at all It would be most interesting, if observers would publish records of failures in the treatment of kala-azar by the antimonyl tartrates, as many seem to believe that the last word about the treatment of the disease has been said in tartar emetic or sodium antimonyl tartrate Unfortunately, this is not so This, coupled with the fact that several cases require more than 4 or 5 months for a cure, demands the need for further advances in the present-day antimony treatment of kala-azar

In the following series of refractory cases, the results obtained from the treatment with the antimonyl tartrates are

first described and the subsequent results obtained after intra venous injection of urea stibamine recorded

(I) Patient named R—History of fever for about six months. When he came to me for treatment his condition was as follows

Fever—temperature 99° to 101°F Spleen felt 6 below the costal margin Blood condition R B C —3 000 000 W B C —3 500 Hb —40 per cent Peripheral blood culture and spleen puncture—positive

Treatment with antimonyl tartrates 6 grammes of sodium antimonyl tartrate and 2.2 grammes of potassium antimonyl tartrate twice a week extending over a period of nearly six months. Altogether 75 injections were given Signs of intolerance began to show themselves after the 30th injection consisting of vomiting diarrhoea severe cramps in the extremities at night during sleep and intense pain in the joints within 24 hours after injection. The treatment was still continued till another 45 injections were given

During treatment patient was also given a course of soamin along with antimonyl tartrates for some time, 2 grammes of soamin having been given in 11 injections. To bring about leucocytosis he was given a trial with 6 injections of T C C O but to no effect

The patient was then sent to Darjeeling where he stayed for two months and a half but without any benefit

Condition of the patient after treatment with antimonyl tartrates Fever—still ranging between 99° to 101°F Spleen felt 5 below the costal margin Blood condition R B C —3 250 000 W B C —4 500 Hb —45 per cent Peripheral blood culture and spleen puncture—positive A refractory case

Subsequent treatment with urea stibamine The injections were given twice a week and the doses were (1) 25 gramme (2) 2 gramme (3) 2 gramme (4) 25 gramme

(5) 25 gramme, (6) 25 gramme, (7) 25 gramme, (8) 25 gramme, (9) 25 gramme Total quantity—2 grammes

The patient began to improve after the second injection the fever stopped after three injections and the condition of the patient was as follows, after the 9th injection

Fever—no rise of temperature Spleen could just be felt below the costal arch Blood condition RBC—5,000,000, WBC—6,250, Hb—70 per cent. Peripheral blood and splenic blood culture and spleen puncture—negative

The patient was under my observation for two months after his treatment was stopped, and when leaving Calcutta he was in excellent health. There was no enlargement of the spleen and the blood condition was R B C —5,000,000, W B C —7,000, Hb —75 per cent. Patient cured

NB—The treatment with urea stibamine was started two months and a half after treatment with sodium antimonyl tartrate was discontinued during which all the antimony present in the patient's system must have been excreted, as I have shown in my paper Chemotherapy of Antimonial Compounds in Kala-azai Infection, Part VIII (Indian Journal of Medical Research, January, 1924) that the average period during which antimony is passed by the kidneys after repeated injections of sodium or potassium antimonyl tartrate is about a month and a half, and therefore the beneficial effects obtained subsequently in the patient could not be due to any residual antimony left in the patient's tissues after the previous treatment

(II) Patient named Miss M—History of fever for five months When she came under my treatment, her condition was as follows Fever—temperature ranging between 99° and 102°F Spleen felt 3½" below the costal margin Blood condition RBC—2,500,000, WBC—3,000, Hb—40 per cent Peripheral blood culture and spleen puncture—positive

Treatment with sodium antimonyl tartrate 4.6 grammes of sodium antimonyl tartrate in 55 injections given twice a week extending over a period of nine months. Signs of intolerance began to show themselves after the 25th injection consisting of palpitation and high fever followed by heavy sweats. The treatment was continued till another 30 injections were given

During the course of treatment there was some improvement in the condition of the patient and the fever stopped for some time but recurred again. The condition of the patient after the course of treatment with sodium antimony! tartrate was as follows. Fever—temperature irregular with periods of apyrexia from time to time. Spleen felt 2¹ b low the costal margin. Blood condition. R.B.C.—3.000.000. W.B.C.—4.000. Hb.—45 per cent. Peripheral blood culture and spleen puncture—positive. A refrictory case.

Subsequent treatment with urea stibamine. Altogether 8 injections were given in doses of 2 gramme each twice a week. The patient began to improve after the third injection there being no rise of temperature and the spleen diminished in size quickly. The condition of the patient one month after cessation of treatment was as follows. Fever—no rise of temperature for nearly two months. Spleen slightly felt below the costal arch. Blood condition R.B.C.—4.000.000. W.B.C.—7.500. Hb.—65 per cent Peripheral and splenic blood culture and spleen puncture—negative. The patient was examined by me again six months after completion of treatment and she was in excellent health. Patient cured.

- (III) Patient J and 24 —The following is the history of his case
- (1) He had two courses of treatment in Dacca with intravenous injection of tartar emetic at an interval of two months between them During the first course he had 1.2 grammes in 14 injections and in the second 8 gramme in

12 injections I-le left the treatment without much benefit, there being no diminution in the height of the fever and in the size of the spleen.

(2) He was admitted into the Berry White Hospital in Dibrugarh after a fortnight. His condition on admission was as follows. Temperature—a double rise ranging between 99° and 103°F. R.B.C.—2,420,000, W.B.C.—2,810, Hb.—50 per cent. Spleen enlarged, extending 3" below the costal arch. Spleen puncture—positive.

Patient was given here a course of treatment with sodium antimony tartrate from 16th November, 1920 to 4th April, 1921, the total dose being 2 grammes. Towards the latter part of the treatment, symptoms of intolerance began to show themselves, such as vomiting, harassing cough and pain in the joints. The last injection was followed by a temporary collapse. There was no improvement in his temperature and the spleen was felt 3" below the costal arch. There was an improvement in the blood condition, the leucocytes numbered 6,000 per c m, but otherwise the condition of the patient was not improved.

(3) He was admitted into my ward on 7th January, 1922, six months after the treatment was stopped. His condition was as follows. Temperature—varying between 99° to 101°F. Spleen 7" below the costal arch. R.B.C.—2,500,000, W.B.C.—2,400, Hb.—30 per cent. Spleen puncture—positive.

The patient was given intravenous injections of tartar emetic twice a week, extending over a period of nearly four months. Altogether 30 injections were given (=3.75 grammes). He was discharged on 18th May, 1923. His condition at the time of discharge was as follows. Freedom from fever after the 18th injection. Spleen $3\frac{1}{2}$ ' below the costal margin. R.B.C.—3,600,000, W.B.C.—4,000, Hb.—52 per cent. Spleen blood culture was positive.

- (4) After a month and a half he was again admitted into the Campbell Hospital in a miserable state L D bodies were found on spleen puncture and the leucocyte count was 2 400 the spleen extending 5 below the costal arch. He was again treated with intravenous injection of sodium and potassium antimonyl tartrate given alternately altogether 2 5 grammes being administered in 30 injections. There was no improvement in the patient s condition. He was kept under observation for nearly two months after which his condition was as follows. Spleen 4 below the costal arch. Temperature ranging between 99° and 100°F. R.B.C.—2 200 000 W.B.C.—3 200 Hb.—48 per cent. Spleen puncture positive. A refrictory case.
 - (5) Subsequent treatment with urea stibamine 3.5 grammes of urea stibamine were given in 15 injections twice a week

Effect of treatment Two months after commencement of treatment Temperature—no rise of temperature for nearly a month RBC—4 500 000 WBC—8 400 Hb—60 per cent Spleen could just be felt below the costal arca Peripheral and splene blood culture and spleen puncture—negative Increase of body weight by one stone

Six months after completion of treatment the patient s condition was as follows. Increase of body weight by one stone. Temperature—no rise of temperature since leaving hospital. R B C -4 200 000. W B C -9 000. Hb -62 per cent. Peripheral blood culture—negative. Spleen not felt below the costal arch. Patient cured.

NB—In this case too the treatment with urea subamine was started when all the antimony from the previous treatment was eliminated from the tissues during the two months when the patient had no antimony treatment

(IV) Patient Mr L—Condition on admission Spleen 7 below the costal arch Temperature—99° to 100°F. Peripheral blood culture and spleen puncture—positive RBC—3,000,000, WBC—2,400, Hb—50 per cent

Treatment with tartar emetic 28 grammes given in 40 injections, extending over a period of six months. Effect of treatment—general condition worse than before, with loss of weight and fever of a low intermittent type. Peripheral blood culture and spleen puncture—positive. Spleen 6" below the costal arch. A resistant case

Subsequent treatment with urea stibamine commenced one month after treatment with tartar emetic was stopped, 1 6 grammes in 9 injections in doses of 1 to 25 gramme being given during one month and a half. Effect of treatment—one month after completion of treatment, general condition improved. Increase of body weight by half stone. Spleen not felt below the costal arch. Peripheral blood culture—negative. Fever stopped after administration of 5 gramme of urea stibamine. Two months after completion of treatment, blood condition was as follows. R.B.C.—4,900,000, W.B.C.—7,800, Hb.—60 per cent. There was increase of body weight by one stone. Patient cured

(V) Patient R—Patient was admitted into hospital after he had 25 injections of 2 per cent solution of sodium antimonyl tartrate and potassium antimonyl tartrate given alternately over a period of three months. Total quantity—1 gramme

Condition on admission Temperature 99° to 100°F Spleen 8" below the costal arch RBC—2,600,000, WBC—1,200, Hb—40 per cent Peripheral blood culture and spleen puncture—positive Patient was suffering from dysentery

Treatment with sodium antimonyl tartrate 4 grammes in 40 injections over a period of four months, given mostly on alternate days

Effect of treatment Temperature—no effect Spleen same as before R B C -2 800 000 W B C -3 200 Hb — 38 per cent Peripheral blood culture and spleen puncture—positive A refractory case

Subsequent treatment with urea stibamine 3 grammes given in 12 injections on alternate days during a period of one month

Effect of treatment Two and a half months after completion of treatment no fever for three months Spleen could not be felt below the costal arch RBC—4 600 000 WBC—7 200 Hb—62 per cent Peripheral blood culture—negative Increase of body weight by one stone Patient cured

(VI) Patient M—Patient was admitted into hospital after he had taken 8 intravenous injections of sodium anti-monyl tartrate Condition Speen 7' below the costal arch Temperature 99° to 101°F RBC—2 300 000 WBC—1 000 Hb—40 per cent Peripheral blood culture and spleen puncture—positive

Treatment with sodium antimonyl taritate 5 grammes injected in 40 injections over a period of five months

Result of blood examination Spleen 5 below the costal arch Fever—slight improvement R B C —3 800 000 W B C —3 600 Hb —48 per cent Peripheral and splenic blood culture—positive A refractory case with slight improvement

Subsequent treatment with urea stibamine $\ 2\ 2\ grammes$ in 9 injections given over a period of two months

Effect of treatment Spleen 2½ below the costal margin Freedom from fever for a month and a half R B C —4 800 000 W B C —6 800 Hb —60 per cent Peripheral and splenic blood culture—negative Spleen puncture—negative Five months after completion of treat ment spleen could not be felt below the costal arch

Increase of body weight by one stone. Peripheral blood culture—negative. R B C —4,800,000, W B.C.—8,000, Hb —62 per cent Patient cured.

(VII) Patient Sarogoo. Condition on admission Temperature—99° to 103°F Spleen 6" below the costal arch RBC—2,600,000, WBC—2,800, Hb—40 per cent Peripheral blood culture and spleen puncture—positive

Treatment with sodium antimonyl tartrate 3 grammes injected in 28 injections extending over a period of four months

Effect of treatment Temperature—no effect Spleen same as before Decrease of body weight by one stone R B C -3,600,000, W B C -3,000, Hb.—40 per cent Peripheral blood culture and spleen puncture—positive A resistant case

Subsequent treatment with urea stibamine 24 grammes injected in 10 injections extending over a period of a month and a half

Effect of treatment: Freedom from fever 14 days after commencement of treatment. Two months after completion of treatment, spleen not felt below the costal arch. Increase of body weight by one stone. R.B.C.—4,300,000, W.B.C.—7,000, Hb.—60 per cent. Peripheral blood culture—negative after 10th injection. Patient cured.

(VIII) Patent N, æt 36, was admitted into my ward in the Campbell Hospital with cancium oris and ædema of the extremities

Condition on admission: Spleen 5½" below the costal arch Peripheral blood culture and spleen puncture—positive RBC—2,400,000, WBC—1,000, Hb—26 per cent Temperature—100° to 103°F

Treatment with sodium antimonyl tartrate: 3 8 grammes given intravenously in 45 injections extending over a period of six months. Effect of treatment. There was no improvement in the general condition, no diminution in the size of

the spleen cedema marked loss of weight by 10 lbs can crum or is—diminished R B C —3 200 000 W B C —2 500 Hb —32 per cent Penipheral blood and spleen blood cul ture—positive Temperature—99° to 101°F Symptoms of intolerance appeared after injection of 2 8 grammes of sodium antimonyl tartrate. A refractory case

Subsequent treatment with urea stibamine 2 8 grammes given in 13 injections extending over a period of two months in doses of 1 to 25 gramme

Effect of treatment General condition—great improvement Increase of body weight by one stone Spleen not felt b low the costal arch RBC—4 100 000 WBC—6 200 Hb—54 per cent Peripheral blood culture—negative Patient cured

(IX) Patient B æt 30 was admitted into my ward in the Campbell Hospital with fever of six months duration and ædema of the extremities

Condition on admission Spleen 7 below the costal arch Spleen puncture and peripheral blood culture—Positive RBC—2 500 000 WBC—2 400 Hb—30 per cent Temperature—99° to 102°F

Treatment with sodium antimonyl tartrate 3.35 grammes of sodium antimonyl tartrate given in 35 injections extending over a period of four months

Effect of treatment Fever came down aftar the 15th injection spleen not reduced in size emaciation with marked cedema RBC—2 200 000 WBC—1 000 Hb—25 per cent Spleen puncture—positive Peripheral and splenic blood culture—positive A refractory case

Subsequent treatment with urea stibamine 2 2 grammes given in 12 injections extending over a period of a month and a half

Effect of treatment General condition improved increase of body weight by 9 lbs peripheral and splenic blood culture—negative R B C —3 600 000 W B C —7 000

Hb - 60 per cent Spleen just felt below the costal arch on deep inspiration Patient cured

A further series of six cases that have resisted treatment with more than four grammes of tartar emetic or sodium antimonyl tartrate are rapidly improving under urea stibamine

Purity of the antimonyl tartrates used. The sodium and potassium antimonyl tartrates used in the above cases were chemically pure and especially made for me for purpose of research.

Possibility of Using Usea Stibamine Intramuscularly

Both Major Shortt and myself are using the compound intramuscularly. The local reaction is generally slight. It is possible that the compound may be advantageously used intramuscularly. A report on this subject will be communicated later on

Urea stibamine, being prepared under perfectly aseptic conditions, is sterile and has been found to be so by being repeatedly tested on culture media. Its solution should not be boiled. The same precautions should be taken in making its solution as in the case of neo-salvarsan.

REMARKS

It will be seen that the duration of treatment and the number of injections required for complete cure of the cases recorded in the present paper were less in this series than in two former series already reported by me (Indian Journal of Medical Research, October, 1922 and October, 1923) This confirms the observations of Major Shortt who also found that the course of treatment required in his cases was much less than what was apparently required in my first two series. In other words, sterilization in these cases must have taken place earlier than I thought and I must

have continued my treatment for a longer period than was really necessary for a complete cure

(B)

Therapcutic Value of Stibamine

Stibamine is the name given by me to p amino phenyl stibinate of sodium (Journal of Tropical Medicine and Hygiene 15th August 1921) It bears the same relation to stibacetin as atoxyl or soamin does to ars acetin. Its chemical and physical properties have already been de cribed and its toxicity tested by me (Indian Journal of Medical Research October 1922) Unless perfectly neutral its solution in water is likely to decompose. The pure alt is fairly stable—In the Indian Medical Galette January 1923 it has been stated by Chopra and Napier that the compound was very easily decomposed. Evidently the substance that they were using was impure

Cases Treated with Stibamine

(I) Patient Rishi Kesh act 30 History of fever—about a year Admitted into my ward at the Campbell Hospital on 25th August 1922 Condition on admission anæmic spleen 5 below the costal arch Temperature—100° to 103°F RBC—2 600 000 WBC—2 600 Hb—40 per cent Peripheral blood culture and spleen puncture—positive

Treatment with stibamine 2.4 grammes were given in 20 injections intravenously in the course of 54 days. Blood culture was positive 26 days after commencement of treatment when 10 injections were given. After completion of treatment the patient improved in weight by 1 stone 2 lbs. Spleen could just be felt below the costal arch. R.B.C.—5 100 000 W.B.C.—8 000 Hb.—62 per cent. Peripheral

and spleen blood culture—negative Spleen puncture—negative Fever stopped after 8 injections Patient cured

(II) Patient Kali, æt 12 History of fever—six months Admitted into my ward at the Campbell Hospital on 22nd August, 1922 Condition on admission Spleen 3½" below the costal arch Temperature—99° to 101°F R B C—3,600,000, W B C—2,400 Hb—50 per cent Peripheral and spleen blood culture—positive Spleen puncture—positive

Treatment with stibamine 12 grammes were given in 18 injections intravenously in course of 48 days Blood culture was positive 24 days after commencement of treatment during which 8 gramme was given in 12 injections. After completion of treatment the patient increased in weight by 1 stone 8 lbs. Spleen could not be felt below the costal arch. R.B. C.—4,200,000, W.B. C.—6,000, Hb.—54 per cent. Peripheral blood culture—negative. Two months after completion of treatment, blood culture—negative. The patient was under my observation eight months after commencement of treatment. Patient cured

(III) Patient D'Cruz, æt 35 years History of fever—a year and a half Admitted into my ward at the Campbell Hospital on 20th August, 1922

Condition on admission RBC—3,100,000, WBC—3,600, Hb—40 per cent Peripheral and spleen blood culture—positive Spleen puncture—positive Spleen 7" below the costal arch Temperature—varying from 98° to 99 6°F

Treatment with stibamine 44 grammes in 30 injections intravenously over a period of 100 days. After the treatment, patient improved in weight by 11 lbs. Fever stopped after 18 injections. Peripheral and spleen blood culture—negative. Spleen puncture—negative. R.B.C.—3,900,000, W.B.C.—6,800, Hb.—50 per cent. The spleen still remained enlarged extending 5" below the costal arch. Three months after completion of treatment spleen was $2\frac{1}{2}$ " below

costal arch No leishmania found on culture of the splenic and peripheral blood Spleen puncture—negative No fever since patient left hospital Patient cured

REMARKS

The above three cases show the curative value of stiba mine No attempt will be made in the present paper to give a comparative estimate of the therapeutic values of stibamine and urea stibamine (Indian Journal of Medical Research October 1922)

OBSERVATIONS

- (1) The curative value of urea subamine and its superiority over the antimonyl tartrates have been established by research conducted by me as well as by Major Shortt in different places
- (2) Refractory cases that resisted treatment with antimonyl tartrates have yielded to urea stibamine
- (3) The short course of treatment and the lesser number of injections required in bringing about a complete cure are striking. Up to now no case of kala azar has been met with which has been resistant to urea stibamine.
- (4) No relapse has been met with among the cases that have undergone complete treatment with urea subamine some of these have been under my observation for nearly two years after completion of treatment
- (5) No relapse has up to now been met with among the resistant cases that have subsequently been cured by urea stibamine. Some of these cases have been under my observation for about a year.
- (6) The therapeutic value of stibamine is recorded in three cases
- (7) The possibility of using urea stibamine intramus cularly is suggested

CHEMOTHERAPY OF ANTIMONIAL COMPOUNDS IN KALA-AZAR INFECTION

[Received for Publication, February 14, 1924]

PART X

FURTHER OBSERVATIONS ON QUANTITATIVE STUDIES IN EXCRETION OF ANTIMONY—THE INFLUENCE OF THE BASIC RADICLE AND OF REPEATED INJECTIONS OF AN ANTIMONYL TARTRATE UPON THE EXCRETION OF ANTIMONY

(A)

THE INFLUENCE OF THE BASIC RADICLE OF AN ANTIMONYL TARTRATE UPON THE EXCRETION OF ANTIMONY

In our last paper on quantitative studies in excretion of antimony, we discussed the excretion of antimony after intravenous injection of tartar emetic. Further observations have since been made to determine the excretion of antimony after injection of the different antimonyl tartrates. In the present paper we shall make a comparative quantitative study of antimony excretion after injection of the following tartrates.

- (1) Tartar emetic
- (2) Sodium antimonyl tartrate
- (3) Lithium antimonyl tartrate
- (4) Ammonium antimonyl tartrate
- (5) Urea antimonyl tartrate,

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Excretion of antimony aft ri jection of tastas emen and ur a atbamine



(1) TARTAR EMETIC

The following are summarised from our previous paper on the excretion of antimony after injection of tartar emetic

- (a) Period of presence of antimony in the urine after a single injection of tartar emetic = 46 days on the average
- (b) The same after repeated injections = 49 days on the average after the last injection
 - (c) Total quantity of antimony excreted in 24 hours —

 1st observation 1/16 6 of the total quantity injected

 2nd observation 1/18 of the total quantity injected

 3rd observation 1/14 of the total quantity injected

Total quantity excreted in seven days -

1st ob ervation $\frac{1}{43}$ of the total quantity injected 2nd observation $\frac{1}{34}$ of the total quantity injected

(2) SODIUM ANTIMONYL TARTRATE

(I) Period of presence of antimony in the unine after a single injection —

1st observation after injection of 2 5 c c of a 2 per cent solution 1 200 hours
2nd observation 1 560
3rd observation 916

Average=51 days

(II) Period of presence of antimony in the urine after repeated injections of sodium antimonyl tartrate —

1st observation —9 c c of a 2 per cent solution given in 4 injections—1 380 hours after last injection

2nd observation —29 c c of a 2 per cent solution given in 13 injections—1,584 hours

3rd observation —32 c c of a 2 per cent solution given in 15 injections—1,056 hours

4th observation —30 c c of a 2 per cent solution given in 14 injections—1,042 hours

Average period = 52 days

Daily Excietion of Antimony in the Urine

1st observation —Injection of 5 c c of a 2 per cent solution (= 0341 grm. Sb)

Day	Amount of urine passed	Amount of antimony passed
1	225 с с	00288 grm
2	375 ,,	002826 ,.
3	525 ,,	0018 ,,
4	720 ,,	00171 .,
5	940 ,,	00117 ,,
6	975 ,,	00099 ,,
7	975 ,,	000936 ,,

Total quantity of antimony passed in 24 hours = $\frac{1}{11.8}$ of the quantity injected

Total quantity passed in seven days = 012312 grm or $\frac{1}{277}$ of the total quantity injected

2nd observation —Injection of 5 c c of a 2 per cent solution (= 0341 grm Sb).

Day	Amount of urine passed	Amount of antimony passed
1	I 250 c c	00324 grm
2	285	002464
3	1 000	0014
4	I 450	0017
5		00115
6	1 100	0006
7	1 625	00068

Total quantity of antimony passed in 24 hours = $\frac{1}{10.5}$ of the total quantity injected

Total quantity passed in seven days = 011234 grm or $\frac{1}{303}$ of the total quantity injected

(3) LITHIUM ANTIMONYL TARTRATE

Period of presence of antimony after a single injection of 5 c c of a 2 per cent solution of lithium initimonyl tartrate

1st observation —478 hours (5 c c of a 2 per cent solution injected)
2nd observation —456 hours ditto

2nd observation —456 hours ditto 3rd observation —312 hours ditto

Average period = 17 days

Daily Exerction of Antimony in the Urine

1st observation — Injection of 5 cc of a 2 per cent solution (= 0406 grm Sb)

\mathbf{Day}	Amount of urine passed	Amount of antimony passed
ī	1 750 c c	006 grm
2	1 900	0028
3		0021
4	2 000	0014
5	1 200	00162
6	1 300	00162
7	1 250	80100

Total quantity of antimony passed in 24 hours = $\frac{1}{6.76}$ of the quantity injected

Total quantity passed in seven days = '01662 grm. or $\frac{1}{2.44}$ of the total quantity injected

2nd observation.—Injection of 5 c c of a 2 per cent solution (= 0406 grm. Sb)

Day	Amount of urine passed	Amount of antimo 13 passed
1	1,000 c c	00445 grm
2	500 ,,	00405
3	320	002754 ,
4	275 .,	00162 ,,
5	425 ,,	00126 .
6	650	0014 ,.
7	550 ,,	000864 ,,

Total quantity of antimony passed in 24 hours = $\frac{1}{91}$ of the quantity injected

Total quantity passed in seven days = 016498 grm or $\frac{1}{2.46}$ of the total quantity injected

3rd observation—Injection of 5 c.c of a 2 per cent solution (= '0406 grm Sb)

Day	Amount of urme passed	Amount of antimony passed
1	1,095 c c	0044 grm
2	950 ,,	004100 ,,
3	830 ,,	003600 ,,
4	2,300 ,,	002146 ,,
5	2,560 ,,	001890 ,,
6	2,650 ,,	001700 ,,
7	2,175 ,,	001379 ,,

Total quantity of antimony passed in 24 hours = $\frac{1}{92}$ of the quantity injected

Total quantity passed in seven days = 019215 grm or $\frac{1}{2.1}$ of the total quantity injected

(4) AMMONIUM ANTIMONYL TARTRATE

(1) Period of presence of antimony in the urine after a single injection of 5 c c of a 2 per cent solution of ammonium antimonyl tartrate —

1st observation	600 hours
2nd observation	576
3rd observation	504
4th observation	480

Average period—23 days approximately

(II) Period of presence of antimony in the unine after repeated injections —

7 injections

888 hours or 37 days

Daily Excretion of Antimony in the Urine

1st observation —Injection of 5 c c of a 2 per cent solution of ammonium antimonyl fartrate (= 0381 grm Sb)

$D_{\mathbf{a}\mathbf{y}}$	Amount of urme passed	Amount of antimony passed
1	1 650 c c	003564 grm
2	1 750	00288
3	2 045	00279
4	2 350	00162
5	1 075	001044
6	3 500	00155
7	1 775	000527

Total quantity of antimony passed in 24 hours = $\frac{1}{10.7}$ of the quantity injected

Total quantity passed in seven days = 013775 grm or $\frac{1}{2.76}$ of the quantity injected

2nd observation —Injection of 5 c c of a 2 per cent solution of ammonium antimonyl tartrate (= 0381 grm Sb)

Day	Amount of urine passed	Amount of antimony passed
l	715 с с	00315 grm
2	575 ,,	002128 ,,
3	675 ,,	00206 ,,
4	320 .,	00161 ,,
5	360 ,,	00129 ,,
6	500 ,,	0012 ,
7	1,375 ,,	0009 ,,

Total quantity of antimony passed in 24 hours = $\frac{1}{12}$ of the quantity injected

Total quantity passed in seven days = 012338 grm or $\frac{1}{31}$ of the quantity injected

3rd observation —Injection of 5 c c of a 2 per cent solution of ammonium antimonyl tartrate (= 0381 grm Sb)

Day	Amount of urine passed	Amount of antimony passed
1	1785 с с	00378 grm
2	1450 ,,	0036 ,
3	1320 ,,	00229 ,,
4	1180 ,,	00129 ,,
5	1760 ,,	001 ,,
6		000788 ,,
7	1230 ,,	000576 ,,

Total quantity of antimony passed in 24 hours = $\frac{1}{10.1}$ of the quantity injected

Total quantity of antimony passed in seven days = 013324 grm or $\frac{1}{2.86}$ of the total quantity injected

(5) UREA ANTIMONYL TARTRATE

Period of presence of antimony in the urine after a single injection of 5 c c of a 2 per cent solution —

1st observation	456 hours
2nd observation	456
3rd observation	384

Average = 18 days

Daily Excretion of Antimony in the Urine

Ist observation —Injection of 5 c c of a 2 per cent solution of urea antimonyl tertrate (= 03175 grm Sb)

Day	Amount of urme pas ed	Amount of antimony passed
i	900 c c	0036 grm
2	1 000	00378
3	780	00165
4	1 200	C0144
5	700	0014
6	1 030	001242
7	l 030	001

Total quantity of antimony passed in 24 hours = $\frac{1}{88}$ of the quantity injected

Total quantity passed in seven days = 14112 grm. or $\frac{1}{22}$ of the total quantity injected

2nd observation —Injection of 5 c c of a 2 per cent solution of urea antimonyl tartrate (= '03175 grm Sb)

Day	Amount of urine passed	Amount of antimony passed
l	525 c c	00330 grm
2	620 ,,	00220 ,,
3	664 ,,	00159 ,,
4	880 ,,	00153 ,,
5	940 ,,	00140 ,,
6	725 ,,	00126 ,.
7		00108 ,,

Total quantity of antimony passed in 24 hours = $\frac{1}{96}$ of the quantity injected

Total quantity of antimony passed in seven days = 01236 grm or $\frac{1}{2.5}$ of the total quantity injected

3rd observation —Injection of 5 c c of a 2 per cent solution of urea antimonyl tartrate (= 03175 grm Sb)

Day	Amount of urine passed	Amount of antimony passed
1	540 c c	00468 grm
2	285 ,,	0021 ,,
3	570 ,,	0024 ,,
4	1,560 ,,	00132 ,,
5		0012 ,,
6	1,560 ,,	00108 ,,
7	1,720 ,,	00081 ,,

Total quantity of antimony passed in 24 hours = $\frac{1}{6.8}$ of the quantity injected

Total quantity of antimony passed in seven days = 01359 grm or $\frac{1}{23}$ of the total quantity injected

CONCLUSIONS

- (1) The proportion of the amount of antimony passed in 24 hours to the amount injected varies with the different antimonyl tartrates and in the following diminishing order
 - (1) Lithium salt-urea salt
 - (11) Sodium salt-ammonium salt
 - (iii) Potassium salt
- (2) The proportion of the amount of antimony passed in seven days after injection of an antimonyl tartrate to the amount injected tends to follow the above order approximately
- (3) The lithium salt is most quickly eliminated next the urea salt next the ammonium salt and lastly the potassium and sodium salts
- (4) Comparing the rate of solubility of the antimonyl tartrates which is in the following order viz (1) lithium salt (2) sodium salt (3) ammonium salt (4) urea salt (5) potassium salt one finds that the amount excreted and the time taken by the urine to be free from antimony does not exactly follow the order of solubility of the salt. In the case of the urea and lithium salts one has to consider their durence action.
- (5) The curves of excretion of the different antimonyl tartrates have been described

(B)

Excretion of Antimony after Repeated Injections of Tartar Emetic

It will be seen from our observations that only a small amount of antimony is excreted by the kidneys after intravenous injection of an antimonyl tartrate during the first twenty-four hours and that even after seven days there is still a fair proportion retained in the system. This fact opens up the question as to whether this retention of antimony by the system continues in the same proportion after repeated doses or whether after the antimony has reached a certain concentration in the system, it is excreted in larger quantities. To determine this we have investigated the excretion of antimony after repeated injections of tartar emetic and the following results were obtained —

[N B—Since most of the antimony injected intravenously is excreted by the kidneys, only a little being passed by the stools, it may be assumed that the amount of antimony present in the tissues on a particular day is represented by the amount of antimony injected minus the amount of antimony excreted by the kidneys]

1st Case

1st injection— $2\frac{1}{2}$ c c of a	2 per cent solution		12	
2nd injection—3 c c of a 2	per cent solution	emetic 7th day	Zinje	ections
Amount of antimony excre	ted on 1st day		00108	grm
Do	2nd ,,		00108	,,
Do	3rd ,,		00099	,
D _o	4th ,,		00054	,,
Dо	5th ,,		000324	,,
$\mathbf{p}_{\mathbf{o}}$	6th ,		000324	,,
D _o	7th ,,		002304	,,
Do.	8ıh ,,		00162	,,
D _o	9th ,,		00153	,,
D ₀	10th ,,		00144	,,
D _o	llth ,		00108	,
р.	12th ,,		00099	,,
Do,	13th ,		000936	,,

- (1) Proportion of antimony excreted to the amount of antimony injected = $\frac{00108}{018}$ or $\frac{1}{167}$ on the 1st day
- (2) Total quantity of antimony excreted up to the 6th day = 004338 grm

Proportion of antimony excreted on the 7th day to the amount of antimony injected on the 7th day plus the amount present in the body due to the previous injection

$$=\frac{002304}{013662+0216}=\frac{1}{153}$$

2nd Case

Ist injection—25 c c of a 2 per cent solution of tartar

2nd injection—3 c e of a 2 per cent solution of tartar emetic on the 5th day from the beginning of obser

3rd injection—4 c c of a 2 per cent solution of tartar emetic on the 8th day from the beginning of obser

injections

Amount of antimony excreted on the 1st day		001296 grm	
Do	2nd	80100	
Do	3rd	0009	
Do	4th	00063	
Do	5th	00282	
Do	6th	0028	
Do	7th	00162	
Do	8th	0038	

- (1) Proportion of antimony excreted to the amount of antimony injected on the first day= $\frac{001296}{018}$ or $\frac{1}{139}$ of the amount injected
- (2) Total quantity of antimony excreted up to the 4th day = 003906 grm

Proportion of antimony excreted on the 5th day to the amount of antimony injected on the 5th day and the amount of antimony present due to the first injection $= \frac{00282}{014094 + 0216} = \frac{1}{12.66}$ which is nearly equal to the proportion on the first occasion

(3) Total quantity of antimony excreted up to the 7th day = 011146 grm

Proportion of antimony excreted on the 8th day to the amount injected on the 8th day and the amount present due to the previous injections = $\frac{0038}{028454 + 0288} = \frac{1}{153}$

It will be seen from the above two observations that the amount of antimony passed by the kidneys was fairly proportional to the amount of antimony present in the tissues in other words the law formulated by us (Indian Journal of Medical Research, January, 1924) that the amount of antimony excreted by the kidneys was fairly proportional to the amount of antimony present held good in those cases in which more than one injection was given

Further observations proved that this law holds good only under certain limits and after several repeated injections of an antimonyl tartrate, a concentration is reached in the tissues at which the amount excreted by the kidneys is greater than what would follow from the above law, so that the concentration of the antimony in the tissues never exceeds this limit. We shall term this limit maximum concentration limit in the tissues. The tendency of the kidneys to throw out antimony from the tissues when it reaches a certain limit in the tissues is no doubt due to the mobility of the threshold of the kidneys, which is lowered when the above concentration is reached. This is illustrated in the following two observations—

(

3rd Case

1st injection-21 c c of 2 per cent solu	tion of tartar em	etic		
2nd injection—3			e 4th da	y
3rd injection-4	do	do	7th	
4th injection—5	do	do		
5th injection—5	do	do	14th	
6th injection—5	do	do	17th	
7.1	do	da	7101	

6th injection—5		do	do	17th
7th injection—5		do	do	21st
Amount of antimony exerci-	ed on the 1st day		00	009 grm
D _o	2nd		00	0063
Do	3rd		0	0065
Do	4th		0	015
Do	ə th		00	0103
Do	6th		CO	18
Do	7th		C	94כם
Do	8th		C	0504
Do	9th		0	029
Do	10th		0	031
D _o	lith		C	0684
Do	12th		00	136
D _o	13th		00	36
Do	14th		00	738
Do	15th		GC	0828
Do	16th		0	0612
Do	17th		0	117
Do	18th		00	i8 7
Do	19th		00)576
Do	20th		00	0648
Do	21st		01	34
Do	22nd ,		00	081

- (I) Proportion of antimony excreted on the 1st day to the amount injected $=\frac{0009}{018} = \frac{1}{20}$ of the amount injected
- $^{(2)}$ The total quantity of antimony excreted up to 3rd day is 00218 grm. Sb

The proportion of antimony excreted on the 4th day to the amount injected on the 4th day plus the amount of antimony already present in the body = $\frac{0015}{(018 - 00218) + 0216}$ $= \frac{1}{249}$

(3) The total quantity of antimony excreted up to 6th day is 00651 grm. Sb

The proportion of antimony excreted on the 7th day to antimony injected on the 7th day plus the amount of antimony already present in the body = $\frac{00594}{(0396 - 00651) \div 0288}$ = $\frac{1}{10.4}$

(4) The total quantity of antimony excreted up to 10th day is 02349 grm. Sb

The proportion of antimony excreted on the 11th day to the antimony injected on the 11th day plus the amount present in the body is $= \frac{00684}{(0684 - 02349) + 036} = \frac{1}{11.8}$

(5) The total quantity of antimony excreted up to 13th day is 03753 grm Sb

The proportion of antimony excreted on the 14th day to the antimony injected on the 13th day plus the amount present in the body is = $\frac{00738}{(1044-03753)+036} = \frac{1}{13.9}$

(6) The total quantity of antimony excreted up to 16th day is 05931 grm Sb

The proportion of antimony excreted on the 17th day to the antimony injected on the 17th day plus the amount present in the body is = $\frac{0117}{(1404 - 05931) + 036} = \frac{1}{10}$

(7) The total quantity of antimony excreted up to 20th day is 09195 grm. Sb

The proportion of antimony excreted on the 21st day to the amount injected on the 21st day plus the amount present

in the body is =
$$\frac{0134}{(1764 - 09195) + 036} = \frac{1}{9}$$

4th Case

Till Case					
1st injection of 25 c c	of 2 per cent solution of	tartar emetic			
2nd injection of 3	do	on the 4th day			
3rd injection of 4	do	do 7th			
4th injection of 5	do	do 11th			
5th injection of 5	do	do 14th			
6th injection of 5	do	do 17th			
7th injection of 5	do	do 21st			
Amount of antimony e	xcreted on the 1st day	001 grm			
Do	2nd	0009			
D _o	3rd	0005			
Do	4th	00198			
Do	5th	00171			
D _o	6th	00144			
Do .	7th	00288			
Do	8th	00216			
Do	9th	0023			
Do	10th	0018			
Do	1 (th	0046			
Do	12th	0029			
Do	13th	0019			
Do	14th	0046			
Do	15th	0063			
Do	16th	00612			
D _o	17th	0123			
Do	18th	00>9			
D _o	19th	0045			
Do	20th	0064ა			
Do	21st	01123			
Do	22nd	0066			

- (1) The proportion of antimony excreted on the 1st day to the amount injected is $\frac{1}{18}$
- (2) The total quantity of antimony excreted up to 3rd day is 0024 grm Sb

The proportion of antimony excreted on the 4th day to the amount injected on the 4th day plus the amount present in the body = $\frac{00198}{(018-0024)+0216} = \frac{1}{187}$

(3) The total quantity of antimony excreted up to 6th day is 00753 grm. Sb

The proportion of antimony excreted on the 7th day to the amount injected on the 7th day plus the amount already present in the body = $\frac{00288}{(0396 - 00753) + 0288} = \frac{1}{211}$

(4) The total quantity of antimony excreted up to 10th day is 01697 grm Sb

The proportion of antimony excreted on the 11th day to the amount injected on the 11th day plus the amount already present in the body = $\frac{0046}{(0684 - 01697) + 036} = \frac{1}{19}$

(5) The total quantity of antimony excreted up to 13th day is 02637 grm Sb

The proportion of antimony excreted on the 14th day to the amount injected on the 14th day plus the amount already present in the body = $\frac{0046}{(1044 - 02637) + 036} = \frac{1}{24.7}$.

(6) The total quantity of antimony excreted up to the 16th day is 04339 grm Sb

The proportion of antimony excreted on the 17th day to the amount injected on the 17th day plus the amount already present in the body = $\frac{0123}{(1404 - 04539) + 036} = \frac{1}{10.8}$

(7) The total quantity of natimony excreted up to the 20th day is 07207 grm Sb

The proportion of antimory exercted on the 21st day to the amount injected on the 21st day plus the amount already present in the body = $\begin{pmatrix} 01123 \\ (1764-07207)+036 \end{pmatrix} = \begin{pmatrix} 11.4 \\ 11.4 \end{pmatrix}$

In the first case (third case in the paper) it will be seen that when the amount of antimony in the tissues reached (1404—05931)+ 036 grm ie 11709 grm the amount of its excretion is suddenly increased. In the second case (fourth case in the paper) when the innount of antimony in the tissues reached (1404—04539)+ 036 or 13101 grm then the same phenomenon was observed. In other words it may be concluded that when the quantity of antimony present in the tissues exceeded these limits, the tendency towards concentration of the drug was prevented by a larger portion of antimony being excreted than that which could be excreted according to the law of excretion already referred to These figures therefore represent the maximum concentration limit of antimony in the tissues of these individuals.

In the paper on the Chemotherapy of Antimonial Compounds in Kala a ar Infection, Part 1'l (Indian Journal of Medical Research Octob r 1923) it was concluded that the results therein pointed to a cumulative action of tartar emetic or at least made the animal susceptible to the next higher dose. It appears from observations in the present paper that no cumulation takes place in the case of the drug above a certinal limit if it is injected at an interval of three or four days.

CONCENTRATION OF ANTIMO 1Y IN DIFFERENT ORGANS

A few observations have been made by us to determine the concentration of antimony in the different organs after 29-767B

- (1) The proportion of antimony excreted on the 1st day to the amount injected is $\frac{1}{18}$
- (2) The total quantity of antimony excreted up to 3rd day is 0024 grm. Sb

The proportion of antimony excreted on the 4th day to the amount injected on the 4th day plus the amount present in the body = $\frac{00198}{(018-0024)+0216} = \frac{1}{18} \tilde{7}$.

(3) The total quantity of antimony excreted up to 6th day is 00753 grm Sb

The proportion of antimony excreted on the 7th day to the amount injected on the 7th day plus the amount already present in the body = $\frac{00288}{(0396 - 00753) + 0288} = \frac{1}{21.1}$

(4) The total quantity of antimony excreted up to 10th day is 01697 grm Sb.

The proportion of antimony excreted on the 11th day to the amount injected on the 11th day plus the amount already present in the body = $\frac{0046}{(.0684 - 01697) + 036} = \frac{1}{19}$

(5) The total quantity of antimony excreted up to 13th day is 02637 grm Sb.

The proportion of antimony excreted on the 14th day to the amount injected on the 14th day plus the amount already present in the body = $\frac{0046}{(1044 - 02637) + 036} = \frac{1}{24.7}$

(6) The total quantity of antimony excreted up to the 16th day is 04339 grm Sb

The proportion of antimony excreted on the 17th day to the amount injected on the 17th day plus the amount already present in the body = $\frac{0123}{(1404 - 04539) + 036} = \frac{1}{10.8}$

(7) The total quantity of antimony excreted up to the 20th day is 07207 grm. Sb

The proportion of antimory excreted on the 21st day to the amount injected on the 21st day plus the amount already present in the body = $\frac{01123}{(1764 - 07207) + 036} = \frac{1}{114}$

In the first case (third case in the paper) it will be seen that when the amount of antimony in the tissues reached (1404—05931)+036 grm ie 11709 grm the amount of its excretion is suddenly increased. In the second case (fourth case in the paper) when the amount of antimony in the tissues reached (1404—04539)+036 or 13101 grm then the same phenomenon was observed. In other words it may be concluded that when the quantity of antimony present in the tissues exceeded these limits, the tendency towards concentration of the drug was prevented by a larger portion of antimony being excreted than that which could be excreted according to the law of excretion already referred to These figures therefore represent the maximum concentration limit of antimony in the tissues of these individuals.

In the paper on the Chemotherapy of Antimonial Compounds in Kala azar Infection Part VI (Indian Journal of Medical Research October 1923) it was concluded that the results therein pointed to a cumulative action of tartar emetic or at least made the animal susceptible to the next higher dose. It appears from observations in the present paper that no cumulation takes place in the case of the drug above a certain limit if it is injected at an interval of three or four days.

CONCENTRATION OF ANTIMONY IN DIFFERENT ORGANS

A few observations have been made by us to determine the concentration of antimony in the different organs after 20-7678 intravenous injection of tartar emetic in lethal and sub-lethal doses. So far it appears that the concentration of antimony is greatest in the liver. The following organs have been examined (1) brain, (2) heart, (3) liver, (4) kidney, (5) the stomach and intestines

Conclusion

There is a maximum concentration limit of antimony in the tissues after repeated injections of tartar cricic, and this is the safeguard against any cumulation of the drug in the tissues when the concentration reaches this point. There is, however, a tendency towards accumulation of the drug in the tissues, after the first two or three injections.

We are much indebted to Dr Bibhuty Bhushan Maity and Dr Siris Chandra Banerjee for co-operating with us in our researches

CHEMOTHERAPY OF ANTIMONIAL COMPOUNDS IN KALA-AZAR INFECTION

IRec d'for P bl c ten June 9 1924]

PART XI

THE VALUE OF UREA STIBAMINE IN THE TREATMENT OF EARLY KALA AZAR

The therapeutic value of urea subamine in the treatment of kala azar has ben established by the observations of Brahmachari Shortt Shortt and Sen Foster and others This antimonial preparation has recently been given an extensive trial in the wards of the Medical College Hospital Calcutta

To Lieut Colonel Barnardo 1 M S Principal and Professor of Medicine and Lieut Colonel McCay IMS Professor of Clinical Medicine at the Calcutta Medical College as well as to Major Shortt 1 M S Special Kala azar Research Officer Shillong and Dr Percy Foster Medical Officer Badlipar Tea Estate lam under deep obligation for their kind courtesy in supplying me with the records of their observations in a series of cases and allowing me to make use of them in this communication

In my paper on the Chemotherapy of Antimonial Com pounds in Kala azar Infection Part IX I have given my observations in the treatment of resistant cases of kala azar with urea stibamine. The present paper is intended to give an account of its value in early cases of the disease From an economic point of view it seems desirable that its value should be assessed in such early cases to determine whether sterilization can be brought about in them more quickly than in chronic cases

By early cases I mean cases in their first attack of fever or in which the duration of the discase did not generally extend more than four months.

The following series of cases show the remarkable rapidity with which sterilization was obtained in them. To give an idea of the observations made by different observers, I have given here some of the cases under the observers mentioned above, in addition to some of mine

I MEDICAL COLLEGE SERIES

(1) Patient P, European male, at 35 years, was admitted into the wards of Lieut-Colonel Barnardo, i.m. s, with history of fever of about a week's duration. He had a previous attack of typhoid-like fever two months before, lasting for about three weeks. Temperature ranged from 102°F to 103°F with double rises and the spleen extended 2" below the costal arch. There was a progressive leucopenia, the leucocytes diminishing from 3,750 to 2,200 in seven days. Blood culture and Widal reaction for B typhosus were negative and no malarial parasites were found.

The diagnosis of kala-azar having been made, patient was given intravenous injection of urea stibamine twice a week, in doses of 0 1 to 0 2 gramme in five injections in the course of a fortnight. Total quantity—0.55 gramme. After three injections temperature came down to normal and the spleen could hardly be felt below the costal arch. Blood culture on N.N.N. medium was negative after the 5th injection. Leucocyte count (1) Before treatment—2,200, (2) after treatment—6,000

(2) Patient M, æt 10, European female, was admitted into the wards of Lieut-Colonel Barnardo, for treatment of remittent fever of three months' duration Temperature

ranged from 102° to 104°F with double rise the spleen extending up to the umbilicus. There was a progressive leu copenia the leucocytes diminishing from 2 600 to 2 200 in seven days time. Blood culture for B typhosus and Widal reaction were negative and no malarial parasites were found.

The diagnosis of kala azar having been made patient was given intravenous injection of urea tibamine in doses of 0.05 to 0.15 gramme in seven injections in course of two weeks. Total quantity—0.8 gramme. The temperature came down to normal after the 4th injection. The blood culture on N N N medium for flagellates was negative after the 6th injection. At the time of discharge which was three weeks after treatment was stopped the spleen and liver could not be felt below the costal arch and blood culture for flagellates was negative. Leucocyte count. (1) Before treatment—2.200. (2) after treatment—6.000

(3) Patient M European male set 7 was admitted into the wards of Lieut Colonel Barnardo t M S for treatment of remittent fever of 18 days duration. Temperature ranged from 100°F to 103°F the spleen extending 2 b low the costal arch. Widal reaction and blood culture for B typhosus were negative. Leucocyte count—2 500. Peripheral blood culture on NNN medium for flagellates was positive.

The diagnosis of I-ala azar having been made the patient was given intravenous injection of urea stibamine in doses of 0.05 to 0.1 gramme in 5 injections in two weeks. Total quantity—0.35 gramme. Temperature came down to normal after two injections the leucocyte count after the treatment was stopped was 6.250. Blood culture for flagellates was negative after 5 injections. Spleen disappeared below the costal arch after 5 injections. At the time of discharge blood culture for flagellates was negative. Leucocyte

- count (1) Before treatment—2,200, (2) after treatment 8,750
- (4) Patient Miss S, æt 33, was admitted into the wards of Lieut-Colonel McCay, I MS, with history of fever of about 10 days' duration. On admission spleen was just palpable below the costal arch. Peripheral blood culture on NNN medium showed the presence of flagellates.

The patient was given intravenous injection of urea stibamine, the doses ranging from 0.05 to 0.15 gramme in 6 injections during 10 days. Total quantity 0.7 gramme Temperature came down to normal after the 3rd injection and the peripheral blood culture for flagellates was negative after the 5th injection. One month after the treatment was stopped the spleen was not palpable below the costal margin Leucocyte count (1) Before treatment—3,200, (2) after treatment—6,000

(5) Patient Miss S, æt 15, was admitted into the wards of Lieut-Colonel McCay, 1 M S, with history of fever of four days' duration. On admission spleen was just palpable below the costal margin, the leucocyte count being 3,800 Peripheral blood culture for flagellates on NNN medium was positive.

The patient was given intravenous injection of urea stibamine, the dose ranging from 0.05 to 0.1 gramme in 4 injections during ten days. Total quantity—0.3 gramme Temperature came down to normal and remained so after 3 injections. The blood culture for flagellates on N.N. medium was negative after 4 injections. Leucocyte count

- (1) Before treatment—3,800, (2) after treatment—6,500
- (6) Patient E. M, æt 15, was admitted into the wards of Lieut-Colonel McCay, implies Ms, with history of fever of three weeks' duration. Patient was in hospital for about a month during which she was found to suffer from remittent fever with double rise and spleen extended ½" below the umbilicus. Peripheral blood was cultured for flagellates.

the result being positive Patient was treated with intra venous injection of urea stibamine twice a week the dose ranging from 0.05 to 0.2 gramme in 8 injections. Total quantity—0.9 gramme. Fever stopped after the 3rd injection and spleen disappeared completely below the costal arch after the 7th injection. Blood culture was found negative after the 8th injection Leucocyte count. (1) Before treatment—3.600. (2) after treatment—6.700.

(7) Patient set 8 H M came under my treatment with history of fever for about two months and an attack of remittent type of fever four months previously. The spleen was three fingers below the costal arch. Peripheral blood culture on NNN medium showed the presence of flagellates and the blood picture was—R B C —3 000 000 W B C —3 500 Hb —60 per cent.

Patient was treated with intravenous injection of urea stibamine the doses ranging from 0.05 gramme to 0.1 gramme the number of injections being six in the course of 12 days. Total quantity—0.4 gramme. After two injections the temperature came down to normal and the spleen could hardly be felt below the costal arch after the 5th injection. Blood culture was negative after the 6th injection. Blood picture. (1) Before treatment—R. B.C.—3.500.000. W.B.C.—3.500. Hb.—60. per cent. (2) after treatment—R.B.C.—4.500.000. W.B.C.—7.500. Hb.—75 per cent.

(8) Patient named D aet 25 came under my treat ment with history of high remittent fever of eight days duration temperature ranging between 103° and 104°F. The spleen could just be felt below the costal arch. Widal reaction for typhoid was positive I in 20. Peripheral blood culture on N N N medium showed flagellates. Patient was treated with intravenous injection of urea stibamine in doses of 0 I gramme each. Altogether 5 injections were given in ten days. Total quantity—0.5 gramme. The fever stopped after the 1st injection. Blood culture on N N N medium.

was negative after the 4th injection Blood picture (1) Before treatment—R B C 3,200,000, W B C —3,600, Hb 46 per cent, (2) after treatment—R B C 4,600,000, W B C —9,000, Hb —70 per cent Two months after completion of treatment, blood culture for flagellates was negative

- (9) Patient named C, æt 25, H M, came under my treatment with history of fever for 10 days. The fever was of high remittent type and the patient was drowsy for four days. Spleen could just be felt below the costal arch Peripheral blood culture on N N N medium showed flagellates. Patient was given intravenous injections of urea stibatine in doses ranging between 0.05 to 0.1 gramme in the course of 14 days. Altogether 0.5 gramme was given in 6 injections. The fever stopped after the 2nd injection and blood culture became negative after the fourth Blood picture. (1) Before treatment—R B C —3,600,000, W B C —1,400, Hb —42 per cent., (2) after treatment—R B C —4,300,000, W B C —6,400, Hb —50 per cent. Blood culture was negative two months after completion of treatment.
- (10) Patient named R, æt 20, H M, came under my treatment with history of high remittent fever for 21 days Spleen could just be felt below the costal arch Peripheral blood culture on N N N medium showed flagellates Patient was given intravenous injection of urea stibamine, the dose ranging from 0.05 to 0.15 gramme. Altogether 6 injections were given in the course of 14 days. Total quantity—0.55 gramme. The fever stopped after the 3rd injection and the blood culture became negative after the 5th injection. Blood picture (1) Before treatment—RBC—3,600,000, WBC—2,400, Hb—36 per cent, (2) after treatment—RBC—4,500,000, WBC—8,200, Hb—60 per cent. Blood culture was negative two months and a half after completion of treatment.

(11) Patient named D came under my treatment with history of fever of three weeks duration Spleen extended three fingers below the costal arch Blood culture for flagellates was positive The patient was given intravenous injection of 0.1 gramme of urea stibamine on 22.5.1924 and on 27 5 1924 Fever stopped after the first injection and spleen could not be felt below the costal arch after the second injection Blood picture (1) Before treatment-RBC-2 300 000 WBC-3 100 Hb-60 per cent (2) after treatment-R B C -5 000 000 W B C -6 000 Hb -- 85 per cent Blood culture-negative on 28 6 1924

II CASES IN PASTEUR INSTITUTE, SHILLONG (MAJOR SHORTT, I M S)

		tr e e		
	Remarks	Patient cured Patient was previously treated with 43 cc of one per cent solution of sodium antimonyl tartrate without benefit	Patient cured	
ht in letion	lncrease in weig posting a let comp treatment	13 <u>1</u> lbs	15 lbs	
ment	Duration of treat	7	20	
bna səru	enoticelar to oVI to innome letot enomedite	0 7 gramme ın 4 ınjec tıons	17 grammes in 8 injec tions	
TMENT	Blood culture	Positive Negative	× Negative	
BEFORE AND AFTER COMPLETION OF TREATMENT WITH UREA STIBAMINE	Blood picture	Hb 60 R B C 4,350,000 W B C 2,600	Hb 60 R B C 3,512,000 W B C 4,000	
	Spleen puncture	Positive	Positive	
	gize of apleen	Up to navel Not felt helow ribs	3 fingers' breadth Just palpable	
la i	Duration of illness to moisancement to moisancement treatment	4½ months	2 months	
	Sex	M 21 yrs	M 15 yrs	

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- Z Z	N & t ve
Hb 65 R B C 5960 000 W B C 7040 000	Hb 60 R B C \$ 100 600 W B C \$ 4000
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III. BADLIPAR SERIES OF CASES (DR PERCY FOSTER)

Sex Age	Duration of illness at com- mencement of treat- ment	BEFORE AND AFTER COM- PLETION OF TREATMENT WITH UREA STIBAMINE		No of injections and total amount of urea	Remarks
		Size of spleen	Spleen puncture	4tibamine	
M 5 yrs	2 months	31 fingers below costal aich 3 fingers	Positive Negative	0 41 gramme in 5 injections	Patient cured Patient vas pre viously treated vith 15 grammes of sodium autimonyl tar trate without any benefit
M 21 3 rs	1 month	31 fingers below costal arch Nil	Positive ×	0 9 gramme in 5	Patient cured
F 6 yrs	1 week	light fingers below costal arch Nil	Positive ×	0 35 giamme in 4 injections	Patient cured
M 28 yrs	3 months	5 fingers below costal arch Nil	Positive ×	2 15 grammes in 10 injections	Patient cured
M 35 yrs	15 days	5 fingers below cos al arch 3 fingers	Positive Negative	2 05 grammes in 10 injections	Patient eured
M 49 yrs	1 month	2 fingers below costal arch Nil	Positive Negative on liver puncture	170 grammes in 8 injections	Patient cured

REMARKS

It is evident from the cases recorded above that urea stibamine cuts short the course of kala azar to a remarkable degree if administered in its early stages. The same con clusions have been arrived at from observations made by different observers in different places. That the drug cuts short the course of the disease in all its stages has already been shown by previous observations. Its beneficial effects in its early stages are however most noteworthy.

It is evident from an economic point of view and in the interests of the miserable sufferers that the use of urea stibamine in the early stages of the disease cannot be over emphasised

I am indebted to my assistants—Sub A sistant Surgeon Bibhuty Bhusan Maity for making flagellate cultures and Sub Assistant Surgeon Sirish Chandra Banerjee for the blood count of my cases described in the present paper

CHEMOTHERAPY OF ANTIMONIAL COMPOUNDS IN KALA-AZAR INFECTION

[Received for Publication June 16 1921]

PART XII

SOME OBSERVATIONS ON THE CONSTITUTION OF UREA STIBAMINE AND STIBAMINE

Urea Stibamine is the name given (Brahmachari ') to the compound formed by treating *p*-aminophenyl-stibinic acid with urea

The constitution of this compound originally suggested was urea salt of p-amino-phenyl-stibinic acid or

$$CO (NH_2)_2 NH_2 C_bH_1 Sb < OH OH$$

In our subsequent research, it was observed that when an aqueous solution of urea stibamine is treated with dilute hydrochloric acid, a precipitate is obtained which does not dissolve in excess of dilute hydrochloric acid. This precipitate is evidently different from p-amino-phenyl-stibinic acid which dissolves in excess of dilute hydrochloric acid. This fact led us to review the constitution of urea stibamine as originally suggested and this constitutes the basis of the present paper.

When urea in aqueous solution is heated above 60°C, cyanic acid is formed as is shown by appearance of a precipitate which is formed on addition of silver nitrate to the solution of urea and which is insoluble in water. The conversion of urea into cyanic acid is best explained by the

constitution of urea as suggested by Werner and the decomposition may be explained by a simple and straightforward change thus —

$$\label{eq:hnco} \text{hn c} \, \langle \overset{\text{NH}_3}{\circ} = \overset{\text{NH}_3}{\underset{\text{HNCO}}{\leftarrow}} \underset{\text{HO CN}}{\longleftarrow} \text{ho cn}$$

When urea is heated with p amino phenyl stibinic acid instead of the urea forming a salt with the acid the cyanic acid formed may combine with its NH radicle giving rise to a carbamino derivative of the acid the NH₁ formed at the same time combining with its acidic portion. The reaction may be expressed as follows—

The decomposition of urea into cyanic acid and ammonia does not take place on heating a solution of urea in excess of methyl alcohol and therefore the formation of the above carbanino derivative should not take place when a solution of urea in methyl alcohol is treated with p amino phenyl stibinic acid even with the aid of heat Experimentally this has been found by us to be the case. Thus when a solution of urea in methyl alcohol is treated with p amino phenyl stibinic acid the latter goes into solution probably giving rise to the formation of a compound having the structure of urea stibamine as originally suggested by one of us. This compound if formed at all must be a very feeble one because when the above solution is treated with

an excess of absolute alcohol, the original p-amino-phenylstibinic acid is precipitated and urea goes into solution

Further experimental evidence in favour of the structure of urea stibamine suggested by us here is given by various chemical properties of urea stibamine and by its quantitative analysis

Purification of urea stibamine.—

2 grammes of urea stibamine prepared in the usual way are dissolved in the least quantity of water and the solution filtered. To the filtrate absolute alcohol is added and the precipitate formed is filtered and washed with absolute alcohol till free from urea. It is then dried in vacuo over fused calcium chloride.

Composition —

Calculated from NH₂ CO NH C, H, SbO OH ONH, H₂O C= 24 7, H=3 23, N=12 3, Sb=35 3 per cent

Found C=24 6, H=4 37 N=12 52, Sb=35 09 per cent

Chemical properties of usea stibamine -

- (1) To an agucous solution of urea stibamine add dilute hydrochloric acid; a precipitate forms insoluble in excess of the acid (distinction from p-amino-phenyl-stibinic acid)
- (2) To an aqueous solution of the substance add alcoholic caustic potash and chloroform, on heating no smell of phenyl-isocyanate is obtained
- (3) One gramme of urea stibamine is dissolved in water and to this strong cooled hydrochloric acid is gradually added till a precipitate that is at first formed is dissolved After cooling the solution 0.2 gramme of sodium nitrite dissolved in water is added to it. The mixture is now treated with alkaline β -naphthol solution. No dye formation takes place

From the above tests it is evident that urea stibamine contains no NH group attached to the benzene nucleus

Preparation of NH CO NH
$$\bigcirc$$
 Sb $\left\langle \begin{array}{c} OH \\ = O \\ ONa \end{array} \right\rangle$ or sodium

salt obtained by the replacement of NH1 from urea stibamine

EXPERIMENTAL

2 grammes of urea stibamine are dissolved in water and to the cooled solution dilute hydrochloric acid is added in excess. The precipitate is washed several times with dilute hydrochloric acid and subsequently with distilled water till it is free from hydrochloric acid. The washed pecipitate is now dissolved in sodium hydroxide solution and the solution subsequently neutralized with dilute acetic acid. The solution is filtered and to the filtrate is added an excess of absolute alcohol till complete precipitation takes place. The precipitate is washed several times with absolute alcohol and then dried in vacuo in a desiccator over fused calcium chloride.

Composition —

Calculated from NH₂ CONH C₆H₄ SbO OH ONa C= 25 54 H = 24 N = 851 per cent

Found C= 25 4 H = 26 N = 84 per cent

Chemical properties -

- (1) It is insoluble in dilute hydrochloric acid
- (2) It does not show the presence of free NH (see properties of urea stibamine)
 - (3) It does not give rise to dye formation (see above)

Both urea stibamine and the above sodium salt on chemical analysis correspond to the formula that they are carbamino derivatives In the case of urea compound, the formula corresponds to the carbamino compound with one molecule of H₂O added to it

STIBAMINE

This is the sodium salt of p-amino-phenyl-stibinic acid and the name was given to the compound by me (Journal of Tropical Medicine and Hygiene, August 15th, 1921)

It appears that in the formation of the sodium salt three molecules of p-amino-phenyl-stibinic acid polymerize with separation of two molecules of water. The sodium salt of the polymerized acid is stable and neutral in its reaction when dissolved in water.

The formula for the polymerized acid will be $(NH_2 C_0H_4 SbO)_3 H_2O_3$

EXPERIMENTAL.

Stibamine is prepared by carefully dissolving p-aminophenyl-stibinic acid in a solution of sodium hydroxide

Composition —

Calculated for (NH₂ C₆H₄ SbO)₃ HO₃Na

C=28 3 per cent, H=2 3 per cent, Sb=47 9 per cent Found C=27 69 per cent, H=2 89 per cent, Sb=47 4 per cent

REMARKS

- 1 In the light of more recent investigations, the constitution of urea stibamine has to be modified from what was originally suggested
- 2 Unlike stibamine it does not undergo polymerization
- 3 Another allied aromatic antimonial compound that does not undergo polymerization is the sodium salt of amido-glycine-p-amino-phenyl-stibinic acid, which was

named (Brahmacharı) stıbglycıne amıde (NH $_2$ CO CH NH C $_6$ H $_1$ SbO OH ONa)

Further work on the reactions between urea and the ortho or para amino aromatic acids is in progress in our labo ratory

My grateful thanks are due to Major Boyd t M S Chemical Examiner to the Government of Bengal for kindly giving my assistant Mr Das facilities for conducting the combustion experiments in his laboratory in connection with the present research

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CHEMOTHERAPY OF ANTIMONIAL COMPOUNDS IN KALA-AZAR INFECTION

[Received for Publication, August 4, 1924]

PART XIII

FURTHER OBSERVATIONS ON DERMAL LEISHMANOID

In the Indian Medical Gazette (1922) and in the Indian Journal of Medical Research (1923), an account was given of the remarkable skin infection which is observed very rarely in patients suffering from kala-azar after they have undergone a course of antimonial treatment and have been apparently cured as far as the infection of the internal organs is concerned

Recently a similar condition was discovered in another patient under almost similar conditions

Previous history —About two years and a half ago, the patient, æt 45, came under the treatment of Brahmachari He presented all the symptoms of kala-azar, eg, double rise of temperature, enlargement of the spleen and liver Blood picture R.B.C.—2,700,000, W.B.C.—2,400, Hb.—34 per cent He was treated with 2 per cent solution of sodium antimonyl tartrate, 26 injections being given in doses of ½ to 5 cc twice a week. When he stopped treatment, his condition was as follows R.B.C.—3,600,000, W.B.C.—4,500, Hb.—46 per cent, spleen 2" below the costal arch, no rise of temperature for nearly a month



Photograph of a patient showing peculiar eruption of the body nine months after completion of the second course of treatment—a case of Dermal Leishmanoid

Six months later he began again to suffer from fever with increase in the size of the spleen and bleeding from the gums Blood picture RBC—2 600 000 WBC—1 200 Hb—36 per cent He was evidently suffering from a relapse as shown by a positive flagellate culture from the peripheral blood He was treated with another course of sodium antimonyl tartrate 20 injections being given when he was apparently cuted

Nine months after completion of the second course of treatment he noticed a peculiar eruption on his body. The lesions appeared at first as two or three nodules over the ear. These subsequently increased in size and other nodules appeared over the skin m different parts of the body. Along with these a number of slightly raised brown patches appeared over the body none of which were anæisthetic (see Plate XXXII). There was no enlargement of the splecn or the liver. Blood condition. R.B.C.—3.900.000. W.B.C.—7.800. Hb.—75 per cent. No flagellates could be developed on culture of peripheral blood from the veins Scrapings from the nodule showed the presence of L.D. bodies and culture of the juice from the nodules gave positive findings.

The histo pathological changes in the skin are described below by Shorit and the location of the parasites with regard to the layers of the skin is indicated

The patient was put on a course of treatment with urea stibamine. Altogether 20 injections of the compound were given intravenously in doses of 1 to 2 grm. After the treatment, the nodules diminished much in size and the patches on the skin diminished to a great extent. After 15 injections had been given the scrapings from the nodules did not show the presence of definite L. D. bodies but a few suspicious looking bodies were found. The patient stopped treatment when he had improved to a great extent. He was accidentally discovered by Brahmachari, three

months after the treatment was stopped, when he appeared to have still more improved but a few nodules and some patches were still present. No opportunity was given on this occasion to examine scrapings from the skin general conclusion is that the patient had benefited to a great extent During the whole period the patient was under treatment for his skin lesions, no signs of internal leishmaniasis were ever observed Apparently, the leishmania affecting the skin of the patient were not absolutely antimonyfast as shown by the improvement in the skin condition under treatment with urea stibamine. The rather slow improvement of the skin makes one think that the parasites were probably more resistant to antimony than ordinary leishmania or were possibly less accessible to the drug than when situated in other tissues. In other words, the case appears to be one of infection of the skin by leishmania rendered somewhat resistant to antimony by previous antimonial treatment of the case

HISTO-PATHOLOGY

BY

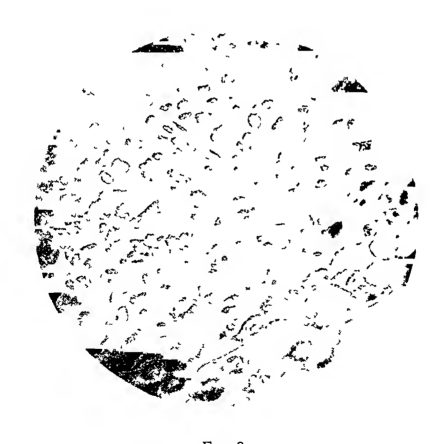
Major H E SHORTT, IMS

Pasteur Institute, Shillong

In 1922, Brahmachari described an interesting condition involving multiple skin lesions in a treated case of kala-azar Since then this condition has been found to be of very rare occurrence, but I had the privilege of seeing a similar, though less marked, case which came under his treatment I am indebted to him for the tissues from which the present description has been prepared.

The lesions, which took the form of papules varying from $\frac{1}{8}''$ to $\frac{1}{2}''$ in diameter, were most numerous in the scrotal region, but were also distributed over the trunk, limbs

[Reprinted from the Indian Journal of Medical Research, Vol XII, No 3, January, 1925] PLATE XXXIII



Fig, 2

Section showing appearance of epidermis and cutis vera in superficial tissue from nodule of a Deimal Leishmanoid case under high power (Vide para 5)

0

and head The description given below applies to a section of one of these nodules cut in a direction perpendicular to the surface of the sLin

APPEARANCES UNDER A LOW POWER (PLATE XXXIII FIG 1)

The first glance shows that there is a profound alteration in the structure of the superficial tissues. This alteration implicates both the epidermis and the cults vera and these will be considered separately.

Epidermis —The structural modification here consists in a uniform attenuation of the epidermis affecting all the recognised layers but most evident in the rete mucosum on account of the relative diminution in number and length of the finger like processes usually associated with this layer in normal skin. While the thickness of the epidermis is thus greatly reduced there is at the same time no tendency for this process to proceed to the extreme degree of ulceration.

Cutis vera—It is in this region that the most striking changes are manifest. The normal condition of dense connective tissue merging gradually into the more open subcutaneous tissue is entirely replaced by what appears to be a dense infiltration of cells forming a deep layer sharply differentiated from the underlying open structure of the subcutaneous tissue. This cellular layer averages in depth about nine times the depth of the epidermis.

APPEARANCES UNDER A HIGH POWER (PLATE XXXIII FIG 2)

Epidermis —Beyond a diminution in the thickness of the layers the minute structure is unaltered

Cutts vera — This appears to be composed of a very primitive connective tissue more or less uniform throughout its extent and formed by rather than infilirated with irregularly shaped cells, many of which have branching

processes The nuclei of the cells are irregularly oval in shape, and stain deeply. The tissue is very vascular and new formation of blood capillaries is actively in progress. The cells composing the walls of the smallest capillaries do not always appear to be distinct from the stroma cells and the impression received from a careful examination of the tissue is that both capillary endothelium and stroma cells are derived from the same primitive undifferentiated cell.

This appears to be the cell which almost invariably contains the intracellular forms of Herpetomonas donovani which are the cause of the condition. Once the primitive cell has become definitely differentiated into a stroma cell or into capillary endothelium, it seems to lose its power of active phagocytosis. The parasitised cells are mainly situated close under the epidermal layer and become fewer in number in the deeper parts of the tissue. Many of these cells, besides containing parasites, enclose abundant pigment granules as the tissue was taken from an Indian patient. The connective tissue underlying the cutis vera is quite normal in appearance.

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CHEMOTHERAPY OF ANTIMONIAL COMPOUNDS IN KALA-AZAR INFECTION

[Rece dfr Puble t n Novemb r 17 1924]

PART XIV

OBSERVATIONS ON A SERIES OF CASES OF KALA AZAR TREATED WITH UREA STIBAMINE DURING A COURSE OF 32 HOURS TO 7 DAYS

In the present paper are described a series of cases of kala azar either under our observation or that of others in which sterilization appears to have been brought about in seven days or less by treatment with urea stibamine. Our grateful thanks are due to Major Shortt 1 M S Lt Col Greig I M S Dr Kundu Dr Percy Foster and Dr Banerjee Badlipar Assam for supplying Brahmachari with notes of their cases By sterilization is meant either negative spleen puncture or negative culture from splenic material or from the peripheral blood together with subsequent disappearance of clinical symptoms of the disease

Cases of Major Shortt 1 M S (Pasteur Institute Shillong)

- (I) Patient et 21 male Fever for four months and a half Spleen puncture—positive Urea stibamine 0.7 grm given in four injections during seven days Sterilization proved by negative microscopic and cultural examination of the splenic juice
- (2) Patient at 16 male Fever for three months Spleen puncture—positive Urea stibanine 0.75 grm given in five injections during seven days Sterilization proved by negative microscopic and cultural examination of the splenic juice

(3) Patient, æt 16, male Fever for eight months Spleen puncture—positive Urea stibamine. 0 65 grm. given in five injections during six days Sterilization proved by negative microscopic and cultural examination of the splenic juice

Cases of Lt -Col Greig, IMS and Di Kundu (Pasteur Institute, Shillong)

(4) Patient No 73—Admitted on 2nd June Spleen puncture on 4th June L D bodies + + + Culture + + + on 10th June Urea stibamine, first injection, 0.2 grm 9th June Second injection, 0.25 grm, 11th June Third injection, 0.25 grm, 13th June Total quantity—0.7 grm

Spleen puncture on the 13th June. on the last day of injection L D bodies—nil Culture—negative after 10 days, that is, on 23rd June

Actual period between the first and third injections—five days Period of illness—four months

(5) Patient No 74—Admitted on 2nd June, 1924 Spleen puncture on 4th June, 1924 L D bodies + + Culture + + + 10th June First injection, urea stibamine, 0.2 grm, 9th June Second injection, 0.25 grm, 11th June Third injection, 0.25 grm, 13th June Total quantity—0.7 grm

Spleen puncture on 13th June, on the last day of injection L D bodies—nil Culture—negative on 23rd June

Actual period between the first and third injections—five days Period of illness—two months

(6) Patient No 164—Admitted on 21st September, 1924 Spleen puncture on 22nd September L D bodies—nil Culture—negative, 1st October First injection, urea stibamine, 0 2 grm, 24th September Second injection, 0 25 grm, 26th September Third injection, 0 25 grm, 26th September Fourth injection, 0 25 grm, 30th September Total quantity—0 95 grm

Spleen puncture on 1st October L D bodies-nil

Culture—negative after ten days, that is, on the 11th October Actual period between first and fourth injections—seven days Period of illness—one year and five months (7) Patient No 171 —Admitted on 25th September Liver puncture 27th September L D bodies—nul Culture—positive on 8th October 1924

First injection of urea stibamine 8th October—0 2 grm Second injection 10th October—0 25 grm I hard injection 12th October—0 25 grm Fourth injection 1-th October—0 25 grm Total quantity 0 95 grm

Liver puncture on 16th October L D bodies-nil Culture-negative 24th October 1924

Actual period between first and tourth injections—seven days Period of illness—about three months

NB-These four cases the notes of which have been very kindly forwarded to Brahmachari are referred to in a joint paper by Lt Col Greig IMS and Dr kundu published in the present number

Cases of Dr Percy Foster and Dr Banerjee (Badlipar Assam)

- (8) Patient act 6 female Fever for seven days Spleen puncture—positive Urea stibamine 0.35 grm given in four injections Stenlization proved by disappearance of the clinical symptoms and subsequent history of the patient
- (9) Patient at 6 female Fever for fifteen days Spleen puncture—positive Urea stibamine 0.5 grm given in four injections during seven days Sterilization proved by negative microscopic examination of the splenie juice and subsequent history of the patient

Cases of Brahmachari and Maity

(10) Patient æt 14 months
Spleen 3° and liver 2° below costal arch Temperature—100
to 102 F Blood culture—positive Urea stibamine 0 05 and 0 1°
gring given intravenously in four days (two impetitions in all) Tem
perature came down to normal after first impetition. Spleen could
not be felt below costal arch after last injection. Actual period
between first and last injections—three days. Peripheral blood
culture—negative after last injection. Period of observation—forty
five days after completion of treatment.

(11) Patient, æt 2 years History of illness—six months Spleen—4½" below costal arch Temperature—101° to 104°F Blood culture—positive WBC—3,000 per cmm Urea stibamine, 0 05 grm, 0 1 grm, and 0 2 grm, given intravenously alternately during five days (three injections in all) Temperature came down to normal after first injection Actual period between first and second injections—two days

Peripheral blood culture—negative after second injection

Period of observation—thirty-six days after completion of treatment, during which spleen disappeared below costal arch and WBC count was 10,000 per c mm

(12) Patient, æt 25 years History of illness—five months Spleen—6½" below costal arch Temperature—apyrexia Blood culture—positive WBC—1,800 per cmm Urea stibamine, 02, 03 and 035 grm, given intravenously on three successive days (three injections in all) Actual period between first and last injections—two days

Peripheral blood culture—negative after last injection

Period of observation—twenty five days after completion of treatment, during which spleen disappeared below costal arch and WBC count was 9,000 per c mm

(13) Patient, æt 18 History of illness—nine months Spleen—4" below costal arch Jaundice—present Temperature—99° to 100°F Blood culture—positive WBC—3,600 per cmm Urea stibamine, 0 2, 0 25 and 0'25 grm, given intravenously on alternate days during five days (three injections in all) Temperature came down to normal after first injection Spleen—disappeared below the costal arch after last injection Actual period between first and last injections—four days

Peripheral blood culture—negative after last injection

Period of observation—twenty-six days after completion of treatment, during which WBC count was 10,400 per c mm and jaundice disappeared

(14) Patient, æt 25 History of illness—three months Spleen —5" below costal arch Temperature—101° to 103°F Patient had commencing cancrum oris Blood culture—positive WBC—1,200 per cmm Urea stibamine, 01 grm, given intravenously on three successive days (three injections in all) Temperature came

down to normal after first injection Actual period between first and last injections—two days

Peripheral blood culture-negative after last injection

Period of observation—fifteen days after completion of treatment during which spleen disappeared below costal arch and W B C count was 5 200 per c mm.

(15) Patient act 35 History of illness—seven months Spleen —5° below costal arch Temperature—100 to 104 F Blood cul ture—positive W B C—1 400 per c mm Urea stibamine 0 l grm given intravenously at (1) 8 a m (2) 12 a m (3) 8 p m on 17th May 1924 and (4) 0 15 grm at 8 a m (5) 0 l grm at 12 a m (6) 4 p m and (7) 8 p m on 18th May 1924 during thirty six hours (seven injections in all) Temperature came down to normal after first injection

Actual period between first and last injections—thirty six hours

Peripheral blood culture—negative after six injections

Period of observation—twenty five days during which spleen disappeared below costal arch and WBC count was 8 000 per c mm

(16) Patient æt 28 History of fever—two months Spleen—4" below costal arch Temperature—101 to 104 F Blood culture—positive W B C—2 400 per c mm Urea stibamine 0.05 gim given intravenously at (1) 6 a m (2) 0.1 grm at 10 a m and (3) 6 p m on 3rd June 1924 (4) 0.1 grm at 0 a m (5) 10 a m (6) 1 p m and (7) 6 p m on 4th June 1924 and (8) 0.1 grm at 12 a m and (9) 4 p m on 5th June 1924 during fifty eight hours (une injections) Tempera ture came down to normal after third injection

Actual period between first and last injections—fifty eight hours Pemberal blood culture—negative after seven injections

Period of observation—six days after which the patient refused to remain under further observation

(17) Patient set 30 History of fever—three months Spleen —6 below costal arch Temperature—99 to 103 F Blood culture—positive WBC—2 800 per c mm Urea stibamine 0 05 grm given intravenously at (1) 6 a m 0 l grm at (2) 10 a m and 0 05 grm at (3) 4 p m on 12th May 1924 and 0 l grm at (4) 6 a m (5) 12 a m (6) 4 p m and (7) 10 p m on 13th May 1924 and 0 l grm at (8) 6 a m (9) 12 a m and 110) 6 p m on 14th May 1924 during sixty hours (ten injections maill) Temperature—normal after mith injection Actual penod between first and last injections—sixty hours

Peripheral blood culture—negative after ninth injection

Period of observation—twelve days, after completion of treatment Spleen—felt 2" below the costal aich during the period WBC count—6,200 per cmm Patient refused to remain under further observation

(18) Patient, æt 7 History of fever—thirty-two days Spleen—21" below costal arch Temperature—99° to 100° F Blood culture—positive WBC—1,800 per c imm Urea stibamine, 0.05 grm, given intravenously at (1) 10 a m and 0.1 grm at (2) 5 p m on 3rd May, 1924, 0.1 grm, at (3) 6 p m on 4th May, 1924, and 0.15 grm, at (4) 4 p m on 5th May, 1924, during fifty-four hours (four injections in all) Temperature—normal after first injection Actual period between first and last injections—fifty-four hours

Peripheral blood culture—negative after last injection

Period of observation—eighteen days, during which spleen disappeared below costal arch and WBC count was 8,800 per e mm

(19) Patient, æt 7 years, History of fever—five months Spleen—extending up to the umbilicus Temperature—98° to 102°F Blood culture—positive WBC—3,400 per emm Urea stibamine, 0 l grm, given intravenously at (1) 8 a m and (2) 4 p m on 20th May, 1924, and 0 l grm at (3) 8 a m and (4) 4 p m on 21st May, 1924, during a period of thirty-two hours (four injections in all) Temperature—normal after second injection Actual period between first and last injections—thirty-two hours

Peripheral blood culture-negative after last injection

Period of observation—thirty-three days after completion of treatment, during which period splcen disappeared below costal arch and WBC count was 11,000 per c mm

(20) Patient, æt 25 History of illness—five months Spleen —5" below costal arch Blood culture—positive WBC—3,100 per cmm Urea stibamine, 0 l grm, injected intravenously twice in the course of five days Temperature came down to normal after the first injection Sterilization proved by complete subsidence of fever, disappearance of the spleen below the costal aich and increase of body weight by 23 lbs in one and a half month WBC count rose up to 6,000 per cmm during this period Blood culture—negative one month and a half after treatment was stopped

(21) Patient æt 9 years History of illness—five months Spleen—2½" below costal arch Liver—slightly enlarged and tender Temperature—99 to 102 F Blood culture—positive WBC—4 200 per cmm One injection of 0.05 gm of urera stibamine and three of 0.1 gm given intravenously in four injections during twelve days Temperature—normal after the first injection Actual period between first and third injections—seven days

Peripheral blood culture—negative after third injection

Penod of observation—five months after completion of treatment and is still under observation Patient has markedly improved in health. No fever during five months. WBC—7500 per cmm one month after completion of treatment.

(22) Patient at 10 Spleen—518 below costal arch Tempe rature—100 to 102 F Blood culture—positive WBC—1800 per c mm Urea stibamine 0 1 6 15 0 2 and 0 25 grm given in six days Temperature came down to normat after second injection Actual period between first and last injections—six days

Peripheral blood culture-negative after last injection

Period of observation—sixty five days during which pleen dis appeared below costal arch and WBC count was 8 000 per c mm

NB—Patient had forty eight injections of sodium antimony tartrate (2.6 grm.) during five months and a half-without any benefit Interval between this treatment and that with urea stibamine—one month and a half

(23) Patient set 25 Spleen—6" below costal arch Tempe rature—99 to 102 F Blood culture—positive WBC—2600 per cmm Urea sthamme 02 025 03 03 and 03 grm given in seven days Temperature came down to normal after first injection Actual period between first and last injections—seven days

Peripheral blood culture—negative after third injection

Period of observation—forty days during which spleen dis appeared below the costal arch and WBC count was 6 800 per c mm

NB—Patient had a course of fifty six injections of sodium antimony tartrate during six months and a half and second course of treatment of twenty four injections of the same and twenty two injections of tartar emetic alternately during seven months linterval between this treatment and that with urea stibamine—four months

(24) Patient, æt 3 Spleen—4½" below costal arch Temperature—100° to 104°F Blood culture—positive WBC—2,600 per cmm Urea stibamine, 0 05 grm 0 1 grm, 0 15 grm, 0 2 grm, and 0 2 grm, given intravenously in five injections during nine days Temperature came down to normal after first injection Actual period between first and fourth injections—six days

Peripheral blood culture—negative after fourth injection

Period of observation—five months Spleen disappeared below costal arch in two months and WBC count was 8,800 per c mm at the end of four months

NB—Patient had forty-five injections of sodium antimony tartrate during five months without any benefit. Interval between this treatment and that with urea stibamine—three months

(25) Patient, æt 25 Spleen—6" below costal arch Temperature—98° to 102°F Blood culture—positive WBC—1,200 per cmm Urea stibamine, 0 25, 0 25, 0 3, 0 3, 0 3, 0 3 and 0 3 grm, given intravenously in seven injections during twelve days Temperature—normal after first injection Actual period between first and fifth injections—seven days

Peripheral blood culture—negative after fifth injection

Period of observation—fifty-five days, during which spleen disappeared below costal arch and WBC count was 7,000 per c mm

NB—Patient had two courses of treatment. During his first course, he had thirty-six injections of sodium antimonyl tartrate (3 grms) and twenty injections of tartar emetic (1 2 grms) during five months and a half. Second course of treatment consisting of fifteen injections of sodium antimonyl tartrate (1 2 grms) and twelve injections of tartar emetic (1 grm) during two months. Altogether, 6 4 grms. Interval between the two courses of treatment and that with urea stibamine—five months.

(26) Patient, æt 20 Spleen—3" below costal arch Temperature—99°F Blood culture—positive WBC—2,800 per cmm

Urea stibamine 0 1 0 1 0 2 and 0 2 grm given intravenously in four injections during five days. Temperature came down to normal after first injection and spleen could not be felt below the costal arch after third injection. Actual period between first and last injections—five days.

Peripheral blood culture—negative after last injection

Period of observation—five months and a half during which spleen disappeared below costal arch and WBC count was 7800 per c mm

NB—Patient had thirty three injections of sodium antimonyl tartrate during four months without any benefit Interval between this treatment and that with urea stibaniune—forty eight days

(27) Patient at 22 Spleen—41" below costal arch Tempera ture—99 to 103 F Blood culture—positive W B C —1 000 per c mm Urea stibamine 0 2 0 2 0 25 0 25 and 0 25 grm given intravenously in five injections during eight days Temperature came down to normal after second injection Actual period between first and third injections—four days

Peripheral blood culture-negative after three injections

Period of observation—three months during which spleen dis appeared below costal arch and WBC count was 6 800 per c mm

NB—Patient had thirty two injections of sodium antimonyl tartrate (22 grms) during four months without any benefit Interval between this treatment and that with urea stibamine—two months and a half

OBSERVATIONS

- (1) The present paper gives a number of cases of kala azar collected from the observations of different observers in which cure was brought about in 32 hours to seven days after treatment with urea stibamine
- (2) The intensive method adopted by us in a few cases of giving multiple injections on the same day so as to main tain a constant high concentration of urea stibamine in the 24-767B

blood has led to the remarkable shortening of the period required for sterilization of the peripheral blood. Though no untoward results were met with in these cases, yet it is too early to state whether the method may be universally advocated.

- (3) The mechanism of response of Leishmania to an antimonial preparation is a very complicated one. While it is universally admitted that urea stibamine brings about sterilization of an infected individual in a much shorter time than tartar emetic or sodium antimonyl tartrate, it is at the same time observed that even with urea stibamine the time required for sterilization is variable. In the present series of cases, this was brought about in seven or less than seven days and in a few cases in thirty-two hours. Though, in some of the cases, striking results were obtained by the intensive method adopted by us in some of our cases, yet there is no doubt that, apart from this, some cases are more quickly amenable to treatment than others. What is the mechanism of this variability? This constitutes an important line of research.
- (4) The percentage of cases very easily amenable to urea stibamine in a series of unselected cases will be most interesting

CHEMOTHERAPY OF ANTIMONIAL COMPOUNDS IN KALA AZAR INFECTION

Re df Pblet Fbu y 18 1925]

PART XV

FURTHER OBSERVATIONS ON CERTAIN DERIVATIVES OF p AMINO PHENYL STIBINIC ACID

The present paper is a continuation of a previous one published in the *Indian Journal of Medical Research*. October 1922 and a second one published in the same Journal October 1924

The preparation of the sodium salt of p amino phenyl stibinic acid which was named stibamine (Brahmachari) has been described in the above mentioned papers

(1) 2 CHLORO I ACETYL AMINO PHENYL 4 SODIUM STIBINATE $C_8H_8O_4NCI$ Sb N_a

For the sake of simplicity we shall name this salt chloro stibacetin

Ten grammes of acetyl p amino phenyl stibinate of sodium are dissolved in 500 c c of water. The solution is cooled with ice and 140 c c of sodium hypochlorite solution (=2 grms of Cl) added to it with vigorous stirring for half an hour. Then dhute acetic acid is added to the above in excess. The whole mixture is vigorously stirred in the cold and filtered after one hour. The precipitate is collected and dissolved in excess of ammonia and filtered. To the filtrate is added dilute acetic acid and the precipitate formed is

filtered and washed with distilled water—It is dissolved in caustic soda solution and neutralized with dilute acetic acid and filtered—After concentrating the filtrate, chloro-stibacetin is precipitated by absolute alcohol and dried in vacuo over fused calcium chloride.

The above compound has the same composition as that of von Heyden's '471'—Stibosan

(2) AMMONIUM SALT OF p-AMINO-PHENYL STIBINIC ACID C₁₉H₂,O₇N₁Sb₃

For the sake of simplicity, we shall name this salt

Preparation

p-amino-phenyl-stibinic acid is dissolved in the least quantity of ammonia in the cold. The acid is reprecipitated by addition of dilute acetic acid. It is filtered and the precipitate is washed several times with distilled water. The precipitate is then dissolved in the least quantity of ammonia and the solution carefully neutralized with dilute acetic acid. It is again filtered and the filtrate is concentrated and then absolute alcohol added to the solution. Ammonium stibamine is precipitated and filtered and subsequently dried in vacuo over fused calcium, chloride.

It may be noted that in the formation of the ammonium salt p-amino-phenyl-stibinic acid polymerizes with the formation of a compound of the following type

$$\begin{array}{c|c}
OH \\
NH_2 & Sb = O \\
O \\
NH_2 & Sb = O \\
O \\
NH_2 & Sb = O \\
ONH_4 & ONH_4
\end{array}$$

The antimony content of the above compound is high and therefore it should be of great therapeutic value in leishmaniasis

The same peculiarity is observed in the case of ammonium stibamine as in the case of stibamine that when a free amino group is present in the benzene nucleus of aromatic amino stibinic acids it polymenizes during salt formation three molecules of the acid polymerizing with separation of two molecules of water

The toxicity of the above compounds will form the subject of another communication

We have observed that glucose has the property of combining with the amino group of p amino phenyl stibinic acid and its derivatives just as it has the property of combining with salvarsan

(3) COMBINATION OF GLUCOSE WITH STIBAMINE

We shall name this compound glucose stibamine

Preparation

Eight grammes of stibamine are dissolved in 40 c c of water containing 6 grms of glucose. The solution is then heated to 60° to 65°C for nearly two hours. It is then filtered and the filtrate concentrated. Absolute alcohol is then added slowly to the filtrate. The precipitate obtained is collected and washed twice with a mixture of alcohol (4 to 1) and subsequently with absolute alcohol. It is subsequently directly over calcium chloride in vacuo.

Glucose stibamine is a yellowish powder very easily soluble in water

(4) COMBINATION OF GLUCOSE WITH AMMONIUM STIBAMINE.

We shall name this compound glucose ammonium stibamine

Preparation

This is prepared in the same way as the above by using ammonium stibamine in place of stibamine. It is less soluble in water than glucose-stibamine.

Detailed observations on these and other allied glucose compounds will be communicated in a subsequent series.

(5) COMBINATION OF GLUCOSE WITH UREA STIBAMINE AND STIBGLYCINE AMIDE

CHEMOTHERAPY OF ANTIMONIAL COMPOUNDS IN KALA-AZAR INFECTION

[Re d f Public t n Febru ry 18 1925]

PART XVI

OBSERVATIONS ON BLOOD CULTURE OF KALA AZAR PATIENTS ON N N N MEDIUM DURING 1922 24

- I COMPARATIVE VALUE OF PERIPHERAL BLOOD CULTURE SPLEEN BLOOD CULTURE AND SPLEEN PUNCTURE IN THE DIAGNOSIS OF KALA AZAR
- II THE PERIOD AT WHICH STERILIZATION OF THE PERIPHERAL BLOOD TAKES PLACE DURING TREATMENT WITH UREA STIBAMINE
- I COMPARATIVE VALUE OF PERIPHERAL BLOOD CULTURE, SPLEEN BLOOD CULTURE AND SPLEEN PUNCTURE IN THE DIAGNOSIS OF KALA AZAR
- (1) In a series of 440 positive cases 426 showed positive peripheral blood culture 1e 97 per cent [These positive cases include (a) 440 cases in which peripheral blood culture was made and found positive in 426 and (b) 190 cases in which splenic culture was made and found positive in all 1
- (2) In a senes of 220 positive cases 186 showed positive spleen puncture results 1 e 845 per cent
- [These positive cases include (a) 160 cases in which peripheral blood culture was made and found positive in 154 cases and (b) 170 cases in which spleen blood culture was made and found positive in all }

II. THE PERIOD AT WHICH STERILIZATION OF THE PERIPHERAL BLOOD TAKES PLACE DURING TREATMENT WITH UREA STIBAMINE

The following table shows the period in which sterilization took place during treatment with urea stibamine

Serial	Age	NUMBER OF IN JECTIONS AND TOTAL AMOUNT OF UREA STIBA- MINE AFTER WHICH BLOOD CULTURE BECAME	Period after com mence- ment of treatment during	Period after com mence ment of treatment at which	Result of blood culture	NUMBER OF TIMES BLOOD CULTURE WAS MADE	
140	No Duration of illness NEGA Period which the amount stibamin give		during blood culture remained positive		during period of observation	During treat ment	During period of observation
1	14 mths 3 mths	2-0 15 grm 72 hrs	76 hrs	80 hrs	Negative (45 days)	4	3
2	2 yrs 6 mths	2-0 15 grm 48 hrs	54 hrs	58 hrs	Negative (36 days)	4	4
3	7 yrs 32 days	4-0 4 grm 53 hrs	62 hrs	70 hrs	Negative (18 days)	9	3
4	7 yrs 5 mths	4-0 35 grm 32 hrs	40 hrs	48 hrs	Negative (28 days)	5	6
5	25 yrs 5 mths	3-0 85 grm 72 hrs	78 hrs	84 hrs	Negative (25 days)	6	6
6	18 yrs 9 mths	3-0 7 grm 144 hrs	148 hrs	152 hrs	Negative (26 days)	10	5
7	25 yrs 3 mths	3-0 3 grm 72 hrs	76 hrs	78 hrs	Negative (15 days)	8	6
8	20 yrs 4 mths	3-0 4 grm 120 hrs	120 hrs	124 hrs	Negative (25 days	6	7
9	35 vrs 9 mths	6-0 65 grm 32 hrs	32 hrs	35 hrs	Negative (28 days)	6	14
10	28 yrs 2 mths	6-06 grm 31 hrs	36 hrs	49 hrs	Negative (6 days)	5	3

_							
Srl N	Ace D ton		mn f f tmnt dig whh	P d aft c m m n e- m nt of t tm t at wh h blood cultur becam eg t	Rult of blood liu d ng p od of bse t n	NUMBER OF TIMES BLOOD CULTURE WAS MADE	
						D 11 g 1 1	Dun g pe d f obse t on
n	30 y 3 mth	9-0 88 grm	52 hr	60 hr	Ng ti (10 dy)	4	2
12	10 y 6 mth	3-072 grm 120 h	12 hr	126 Ъ	Neg t (65 d y)	6	6
13	25 y 6 mth	3-0 75 grm 96 h	106 Ъ	120 h	N g t (40 d y 1	8	6
14	3 y 4 mth	4-0 35 grm	148 hr	155 hrs	Neg (1150 d y)	9	17
15	24 y 6 mth	5-14g m 108 h	168 hr	172 h	Ngt (51 dy)	13	18
16	20 y 4 mth	120 h	168 Ъ	192 Ъ	Ng((165 dy)	15	13
17	22 y 2 mth	5-1 15 g m	204 hr	216 Ъ	N g t (90 d y)	14	16
18	6 mth	6-0 6 g m 240 h s	288 hr	312 h	Ngtv (65 dy)	11	16
19	9 y 6 mth	3-0 15 g m	216 h	240 hr	Ngt (90 dy)	7	4
20	9 mth	240 h	264 h	288 h	Ngt (34 dy)	12	6
21	30 y 6 mth	- 6-08g m 197 h s	226 h	249 h	N g t (36 d y)	13	12
22	mth	230 h s	234 h	240 h	N g t (26 d y)	12	8
23	26 y 11 mth	6-08g m 192 h	204 h	216 h	Ngt (65 dy)	10	9
24	20 y 5 mth	5-08g m	180 h	192 h	N g t t29 d y \	8	12

Serial No	AGE Duration of illness	OF UREA STIBA- MINE AFTER WHICH BLOOD URATION CULTURE BECAME	Period after commence- ment of treatment during which blood culture remained positive	Period after com mence- ment of treatment at which	Result of blood cul- ture during	Number of times blood culture was made	
				blood culture became negative	period of observation	During treat- ment	During period of observation
25	21 yrs 3 mths	5-0 8 grm 14 days	14 days	15 days	Negative (30 days)	9	10
26	8 yrs	4-0 55 grm 264 hrs	300 hrs	312 hrs	Negative (60 days)	21	14
27	11 yrs 6 mths	4-08 grm 264 hrs	276 hrs	288 hrs	Negative (92 days)	14	23
28	18 yrs 5 mths	3-0 4 grm 168 hrs	192 hrs	216 hrs	Negative (75 days)	9	12
29	9 yrs 4 mths	4-0 4 grm 264 hrs	312 hrs	336 hrs	Negative (65 days)	13	10
30	28 yrs 4 mths	9-18 grms 360 hrs	384 hrs	408 hrs	Negative (60 days)	17	12
31	25 yrs 5 mths	5-1 grm 192 hrs	192 hrs	216 hrs	Negative (72 days)	8	12
32	10 yrs 6 mths	6-06 grm 192 hrs	204 hrs	216 hrs	Negative (90 days)	12	14
33	25 yrs 5 mths	6-12 grms 296 hrs	308 hrs	320 hrs	Negative (30 days)	15	14
34	8 yrs	4-0 8 grm 144 hrs	216 hrs	240 hrs	Negative (60 days)	10	10
35	16 yrs 3 mths	5-08 grm 168 hrs	180 hrs	192 hrs	Negative (40 days)	10	9
36	26 yrs 5 mths	3-0 85 grm 17 days	17 days and	18 days	Negative (45 days)	15	11
37	42 yrs 9 mths	4-1 15 grms 28 days	12 hrs 32 days	34 days	Negative (56 days	14	12
		<u> </u>	<u> </u>	1	<u>' </u>	-	

Making a summary of the above cases we can dittem into the following groups —	vide
(I) Number of cases in which sterilization took place in from 32 hours to 84 hours (less than four days)	9
(2) Number of cases in which sterilization took place in from above 84 hours to 144 hours (6 days)	3
(3) Number of cases in which sterilization took place in from above 144 hours to 240 hours (10 days)	15
(4) Number of cases in which sterilization took place in above 10 days to 16 days	7
(5) Number of cases in which sterilization took place in more than 16 days	2
(6) One case required 34 days for sterilization	Ī

From the above we conclude-

26 out of 37 cases 1 e 73 per cent of the cases were sterilized within 10 days after commencement of treatment with urea stibamine

CHEMOTHERAPHY OF ANTIMONIAL COMPOUNDS IN KALA-AZAR INFECTION

[Received for Publication, March 19, 1925]

PART XVII

FURTHER DETAILS OF THE PREPARATION OF UREA STIBAMINE

In the Indian Journal of Medical Research (October, 1922), I described the method of preparing urea stibamine. The details of preparing acetyl-p-amino-phenyl-stibinic acid and p-amino-phenyl-stibinic acid which are two substances required in the preparation of urea stibamine, not having been described in the above paper, are described below —

(1) Preparation of acetyl-p-amino-phenyl-stibinic acid

One gram-molecule of para-amino-acetanilide is added to well-cooled sulphuric acid (1.5 gram-molecule) in one litre of water. The mixture is then diazotised with a cooled solution of sodium nitrite (one gram-molecule) in water. The solution is then added to a solution of sodium antimonite which is rapidly cooled to 0°C. [The sodium antimonite solution is prepared as follows 600 grammes of sodium hydroxide are dissolved in 3 litres of water. The solution is added to aqueous antimony trichloride prepared by dissolving antimony trioxide (5 gram-molecule) in 764 grammes of hydrochloric acid (D 1·123).] After the reaction is complete, the solution is almost neutralised with dilute sulphuric acid and the remainder of the caustic alkali is removed by saturating the solution with carbon dioxide. The solution

is now filtered and the filtrate is saturated with sodium chlo ride when acetyl p amino phenyl stibinate of sodium is precipitated. The precipitate is then dissolved in water and again saturated with carbon dioxide. The solution is filtered and the filtrate treated with dilute hydrochloric acid which precipitates acetyl p amino phenyl stibinic acid.

(2) Preparation of p amino phenyl stibinic ocid

One part of acetyl p amino phenyl stibinic acid is heated for some hours with 10 parts of 5 per cent aqueous sodium hydroxide for some hours until the dilute sample gives with dilute hydrochloric acid a precipitate which dissolves in excess of the acid. To the cooled solution is added dilute acetic acid when p amino phenyl stibinic acid is precipitated.

The p amino plienyl stibinic acid is then suspended in water and urea added to the suspension till the whole of the acid is almost dissolved. The solution is facilitated by gentle heating. It is then filtered and the filtrate concentrated on the water bath. The concentrated solution after cooling is mixed with excess of alcohol which precipitates urea stibamine. The precipitate obtained is further washed with alcohol to free it from any uncombined urea It is subsequently dried over a porous plate.

The constitution of urea stibamine has been fully discussed by myself and my chemist Mr Judhisthir Das in the Indian Journal of Medical Research (October 1924)

CHEMOTHERAPY OF ANTIMONIAL COMPOUNDS IN KALA-AZAR INFECTION

[Received for Publication, September 23, 1925]

PART XVIII

FURTHER OBSERVATIONS ON CERTAIN DERIVATIVES OF p-AMINO PHENYL-STIBINIC ACID (Continued)

This paper is a continuation of the paper entitled Chemotherapy of Antimonial Compounds in Kala-azai Infection, Part XV, published in the Indian Journal of Medical Research, July, 1925

(6) GLUCOSE COMPOUNDS WITH p-AMINO-PHENYL-STIBINIC ACID AND ITS DERIVATIVES

These, already referred to in our paper (July, 1925), consist of glucose compounds with (1) stibamine, '(2) ammonium stibamine, (3) urea stibamine, and (4) stibglycine amide

(7) CONDENSATION OF DICHLORO-ACETAMIDE WITH p-AMINO-PHENYL-STIBINIC ACID

Experimental

Four grammes of p-amino-phenyl-stibinic acid are treated with a watery solution of caustic soda till it dis-

solves The solution is made slightly alkaline with a slight excess of caustic soda. To the solution one gramme of dichloro acetamide is added and the resulting mixture is heated at 60° to 70°C the solution being kept slightly alkaline by addition of small quantities of caustic soda from time to time. It is then filtered and the filtrate cooled in ice. Dilute hydrochloric acid is now added in excess to the solution and the precipitate obtained is washed twice in dilute hydrochloric acid and subsequently five times in distilled water. The precipitate is then dissolved in caustic soda and the solution subsequently made neutral. This solution is filtered and concentrated and to the concentrated solution absolute alcohol is added in excess. The precipitate obtained is filtered and washed in absolute alcohol and dried over calcium chloride in a vacuum desiccator.

(8) THE SODIUM SALT OBTAINED BY THE REPLACEMENT OF NH, FROM UREA STIBAMINE BY N_{Δ}

This has already been described in Indian Journal of Medical Research October 1924. It is sodium carbamino p stibanilate. Its glucose compound has also been prepared

The glucose compounds described in the present paper are prepared in the same way as the glucose compounds with stibamine and ammonium stibamine (Indian Journal of Medical Research July 1925)

Therapeutically the glucose compounds are weaker than the corresponding antimony compounds from which they are derived

CHEMOTHERAPY OF ANTIMONIAL COMPOUNDS IN KALA-AZAR INFECTION

(New Series)

PART I

THERAPEUTIC VALUE OF N-PHENYL-GLYCINE-AMIDE-p-STIBINATE OF SODIUM

(A PRELIMINARY NOTE)

The following is a note of the first series of cases of kalaazar treated with N-phenyl-glycine-amide-p-stibinate of sodium by Brahmachari and co-worker

1 Patient named D, æt 16, was admitted into Brahmachari's ward, with history of fever for 6 months. On admission, his spleen extended five inches below the costal arch. Blood culture on NNN medium—positive flagellate culture. Patient was treated with intravenous injection of the above compound. Altogether 20 injections were given. As a result of treatment the fever completely stopped, the spleen could not be felt below the costal arch, and at the time of discharge blood culture on NNN medium was negative.

Dose— 05 to 20 gramme Increase of body weight after treatment—1st 1lb

Result of Blood Examination—

(1) R B C —1,800,000, W B C —2,500, Hb —35% on 11-2-25 before treatment

- (2) R B C -4 500 000, W B C -8 400 Hb -85% on 20 6 25 after treatment
- 2 Patient named J act 10 was admitted into Brahmachari s ward on 17 11 24 with history of fever for five months. On admission his spleen extended four inches and a quarter below the costal arch. Patient was treated with intravenous injection of the above compound. Altogether 15 mjections were given. The fever stopped after the 3rd injection.

Dose—05 to 20 gramme Increase in body weight after treatment—1 stone

Result of Blood Examination-

- (I) R B C -2 500 000 W B C -3 500 Hb -45% on 19 11 24 before treatment
- (2) R B C -4 400 000 W B C -9 600 Hb -45% on 27 3 25 after treatment
- 3 Patient named S set 14 was admitted into Brahmachari s ward on 3 5 25 with history of fever for three months. On admission his spleen extended 21 inches and the liver 1 inch below the costal arch. Patient was treated with intravenous injection of the above compound Altogether 12 injections were given. Fever stopped after the 4th injection.

Dose- 00 to 20 gramme

Result of Blood Examination-

- (1) R B C $-2\,000\,000\,$ W B C $-1\,800\,$ Hb $-40\,\%$ on 20 6 25 before treatment
- (2) R B C -4 200 000 W B C -7 100 Hb -70% on 12 8 25 after treatment
- 4 Patient named J set 10 was admitted into Biahmachari s ward on 2 3 1925 with history of fever for four months On admission his spleen extended up to 26-767B

the umbilicus and the liver was just palpable below the costal arch Patient was treated with intravenous injection of the above compound Altogether 10 injections were given The temperature came down to normal after the 4th injection At the time of discharge, the spleen and the liver could not be felt below the costal arch

Dose-'05 to 20 gramme

Result of Blood Examination—

- (1) R B C -2,500,000, W B C -2,500, Hb.-45% on 8-3-25 before treatment
- (2) R B C -4,200,000, W B C -6,800, Hb -80% on 10-6-25 after treatment
- 5. Patient named S, æt 13, was admitted into Brahmachari's ward on 17-3-1925, with history of fever for twelve months. On admission, his spleen extended 2 inches below the costal arch. The patient was treated with intravenous injection of the above compound. Altogether 20 injections were given. The temperature came down to normal after the 4th injection. At the time of discharge the spleen could not be felt below the costal arch.

Dose — 05 to 20 gramme Increase in body weight after treatment—1 st 1 lb

Result of Blood Examination—

- (1) R B C ---3,500,000, W B C ---3,100, Hb ---65% on 18-3-25 before treatment
- (2) R B C —4,000,000, W B C —5,100, Hb —75% on 10-5-25 after treatment
- 6 Patient named M., æt., 20, was admitted into Brahmachari's ward on 25-5-25, with history of fever for two years. On admission, his spleen extended 6 inches below the costal arch. Patient was treated with intravenous injection of the above compound. Altogether 15 injections

were given The temperature came down to normal after the 5th injection. At the time of discharge the spleen was felt 2 inches below the costal arch

Dose-05 to 20 gramme

Result of Blood Examination—

- (1) R B C -- 2 500 000 W B C -- 2 100 Hb -- 45 to on 27 5 25 before treatment
- (2) R B C -- 3 200 000 W B C -- 6 500 Hb -- 70% on 17 8 25 after treatment

7 Patient named S æt 21 was admitted into Brahmachari s ward on 28 2 1925 On admission both his spleen and the liver extended 4 inches below the costal arch Patient was treated with intravenous injection of the above compound. Altogether 14 injections were given The temperature came down to normal after the 5th injection. At the time of discharge the spleen was felt I inch below the costal arch.

Dose-05 to 20 gramme

Result of Blood Examination-

- (1) R B C 2 820 000 W B C 1 800 Hb 30 / on 31 3 25 before treatment
- (2) R B C -- 3 000 000 W B C -- 5 000 Hb -- 65% on 29 5 25 after treatment
- 8 Patient named B set 12 was admitted into Brahmachari s ward on 23 3 1925. On admission his spleen extended 8 inches and his liver 2 inches below the costal arch. Patient was jaundiced. He was treat ed with intravenous injection of the above compound Altogether 10 injections were given. The temperature came down to normal after the 6th injection. At the time of discharge the spleen could not be felt below the costal arch.

Dose — 05 to 20 gramme Increase of weight after treat ment—1 st 7 lbs

Result of Blood Examination—

- (1) R B C -2,500,000, W B. C -2,800, Hb.-20% 26-3-25 before treatment
- (2) R B C -4,500,000, W. B C.-6,200, Hb.-80% 10-6-25 after treatment.

REMARKS

This paper gives records of preliminary observations hade by Brahmachari on the treatment of kala-azar ith N-phenyl-glycine-amide-p-stibinate of sodium. In this aper it is not possible to make any comparison of its value with that of other aromatic antimonials. Further investigation in progress.

CHEMOTHERAPY OF ANTIMONIAL COMPOUNDS IN KALA-AZAR INFECTION

(New Series)

PART II

THE TOXICITY AND THERAPEUTIC VALUE OF OLD SAMPLES OF UREA STIBAMINE

Samples of urea stibamine kept for indefinite periods in sealed ampoules under ordinary conditions in India have been tested to determine whether they differed in their toxicity and therapeutic value when compared with fresh samples. The samples tested were those that had been kept in sealed ampoules from the beginning of the year 1922. Their toxicity was tested in January 1926.

Lethal Effects Produced from the Administration of a 2 per cent Solution of Urea Stibamine to Guinea pigs by Intramuscular Injection

Toxicity of fresh samples of urea stibamine

D ng m pkl of bdywght	N mb f	Nmb f gu pg dd	
70 65 60 50 45 45 35	4 3 4 2 4	4 2 2 1	Mınımum lethal dose
45 35	6	l mt	Maximum tolerated dose

Toxicity	of old	samples	of	шеа	stıbamıne
----------	--------	---------	----	-----	-----------

Dose in gram per kilo of body weight	Number of guinea pigs used	Number of guinea pigs died	
70 65 60 45	6 3 3 3	6 2 2 !	Mınımum lethal dose
40 35	6	nıl	Maximum tolerated dose

It will be seen that there is no difference between old and new samples of urea stibamine in their toxicity as tested on guinea-pigs

Physical and Chemical Properties

No difference was observed in the physical and chemical properties between old and new samples of urea stibamine kept in sealed ampoules

Therapeutic Properties

No difference was observed in the therapeutic value of old samples of urea stibamine as compared with new ones. No untoward symptoms were met with during the course of treatment with these samples. Their therapeutic value was tested in three cases. One of them was a case suffering from acute kala-azar with high fever and delirium, and other severe constitutional symptoms. Blood culture for flagellates was positive. The temperature came down to normal after three injections of an old sample of urea stibamine kept in sealed tubes since 1922. After 8 injections were given blood culture was found to be negative and then the injections were stopped. The second case recovered after 10 and the third after 12 injections of the same samples of urea stibamine.

CHEMOTHERAPY OF ANTIMONIAL COMPOUNDS IN KALA AZAR INFECTION

(New Series)

PART III

DERMAL LEISHMANOID WITH POSITIVE FLAGELLATE
CULTURE FROM THE PERIPHERAL BLOOD

Under the name of dermal leishmanoid a rare form of skin lesion was originally described by Brahmachari in 1923 which was characterised by a multiple infection of the skin by Leishmania donovani in individuals who had previously suffered from kala azar (visceral leishmaniasis) and were subsequently cured by a course of antimonal treatment. This observation has since been confirmed (Brahmachari Megaw Knowles Acton and others)

Very rarely cases may be observed in which similar skin lesions due to leishmania donovani infection may be found developing in an imperfectly cured case of kala azar and in which the skin lesions more or less persist after sterilization of the internal organs against leishmania by a course of antimonial treatment

The following is one such extremely rare case and is therefore of exceptional interest

Patient named G A Mahomedan, was at first treated for kala azar by a course of treatment with sodium antimonyl tartrate Six months later he was admitted into hospital with nodular eruptions all over the body. He stayed in hospital for nearly three weeks during which he was treated with 5 injections of Von Heyden s 471 (stibosan) without

any improvement He had also had treatment with galvanic current exposures in the Calcutta School of Tropical Medicine without any benefit Patient came under our observation on 23rd July, 1926, about six months after he had left treatment

Condition at the time of our first observation—Patient had very well marked nodular eruptions over his face, the trunk and the extremities (see Plate) Besides these, there were patches of a-pigmentation over the body especially the trunk He was suffering from fever with enlargement of the spleen

Blood examination on 28th July, 1926, before treatment with urea stibamine (Brahmachari) was as follows —

R B C4,000,000	Leucocyte count—	
W B $C - 2,700$	Polymorphonuclears	61%
Hb —75%	Mononuclears	18%
	Lymphocytes	17%
	Eosinophiles	4%

No malarial parasites Culture from the peripheral blood on Kligglers' media gave positive flagellate culture Wasserman reaction—negative. The nodules in the skin showed Leishmania-donovani on smear and culture

Patient was put on a course of treatment with urea stibamine (Brahmachari) from 15th August, 1926. The spleen quickly diminished in size and the fever stopped. The skin eruptions seemed to be very resistant to treatment, though they slowly diminished from the trunk and extremities after a course of treatment of bi-weekly intravenous injections with urea stibamine extending over a period of five months.

Blood examination on 13th September, 1926, after treatment with urea stibamine (Brahmachari) —

R B C —4,500,000	Leucocyte count	
W B C6,600	Polymorphonuclears	61%
Hb —75%	Mononuclears	18%
	Lymphocytes	19%
	Eosmophiles	2%

Blood culture for flagellates—negative

ORSERVATIONS

The present case is of exceptional interest for the following reasons —

- (1) The patient developed dermal leishmanoid at a stage when he was not completely cured of kala azar as shown by positive flagellate culture from the peripheral blood leucopenia with fever and enlargement of the splean. Though positive flagellate culture from the peripheral blood could be due to the blood having been accidentally infected from the skin at the time of drawing the blood yet the clinical picture of the case at the time when it came under observation showed that the patient was not cured of kala azar.
- (2) He was subsequently cured of Irala azar after a course of treatment with uren stibamine (Brahma chari)
- (3) The skin eruptions are very resistant to treatment with antimony as they seemed to be only very slowly yielding to treatment with urea stibamine

The case appears to have been one of incompletely cured kala azar at the time of our first observation and was met with at a stage when visceral leishmaniasis was passing into dermal leishmanoid lit is therefore the first of its kind as up to now all cases of dermal leishmanoid that have been recorded were found to have developed in kala azar patients after a definite period of recovery from visceral leishmaniasis

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CHEMOTHERAPY OF ANTIMONIAL COMPOUNDS IN KALA-AZAR INFECTION

(New Series)

PART IV

A RARE CASE OF DERMAL LEISHMANOID +

After attention was drawn by Brahmachari to skin eruptions due to infection with leishmania-donovani, Acton in collaboration with Napier has described the following stages of this disease —

- (1) An early de-pigmented stage
- (2) A later nodular stage
- (3) A xanthoma-type of the disease in which there is a tendency towards fibrous tissue formation, and constriction of venules and subsequent dilatation

It is stated by these authors that the nodules appear in the place of de-pigmented patches

Generally speaking, cases of dermal leishmanoid come for treatment when both the nodules and de-pigmented patches are present in the skin. The present case is therefore of much interest, as it showed an erythematous patch over the face and a few nodules over it with complete

^{*} Rend at a meeting of the Medical Section of the Asiatic Society of Bengal, April 16, 1928

absence of de pigmentation in any part of the body confirm ing the view of Brahmachari that the disease may first show itself in the form of patches of erythema over the skin before the appearance of de pigmentation or nodules

History—

About three years ngo the patient had an attack of kala azar. He was treated with about thirty injections of sodium antimonyl tartrate and was apparently cured. About a year ago he noticed a small patch over the bridge of the nose which gradually extended and a sumed the present size. A few papules appeared subsequently over the erythematous area.

Present Condition-

Patient is a healthy individual. There is no enlargement of spleen or liver. There is no fever. Blood culture is negative. There is no anæsthesia over the crythematous patch nor over any other part of the body nor is there any thickening of the nerves. A careful examination of the scrapings from the face showed the presence of a few Leisliman Donovan bodies after a prolonged examination. No lepra bacilli were found from these scrapings.

The case is of much interest as such cases rarely come under observation. Generally speaking cases that present themselves for treatment show definite de pigmented areas with or without nodules either inside them or in independent foci in the skin.

STUDIES IN KALA-AZAR AND CHEMOTHERAPY OF ANTIMONY

- I SUBSEQUENT HISTORY OF THE FIRST RECORDED CASE OF DERMAL LEISHMANOID
- II SUBSEQUENT HISTORY OF A CASE OF DERMAL LEISHMANOID ORIGINALLY CONSIDERED TO HAVE BEEN REFRACTORY TO TREATMENT
- I SUBSEQUENT HISTORY OF THE FIRST RECORDED CASE OF DERMAL LEISHMANOID

The first recorded case of dermal leishmanoid was published by Brahmachari in the Indian Medical Gazette in April, 1922, and described in greater detail in the Indian Journal of Medical Research in April, 1923 The case remained under treatment for irregular periods, and was afterwards lost sight of The following subsequent history has been obtained from one of his relations. The de-pigmented patches increased in size in course of time, and in some places gave rise to typical leucoderma-like areas The nodules ulcerated, giving rise to sanious foul-smelling discharge They subsequently diminished in size, and the ulcers healed up The patient became more and more wasted, and died last February of an attack of dysentery. This is perhaps the first case of dermal leishmanoid in which ulceration ultimately took place in the affected areas of the skin after a long period As, however, the case was not seen by us in the stage of ulceration, the

authenticity of the formation of ulcers must remain a doubt ful point

II SUBSEQUENT HISTORY OF A CASE OF DERMAL LEISHMANOID ORIGINALLY CONSIDERED TO HAVE BEEN REFRACTORY TO TREATMENT

This case came under the observation of Brahmachan in July 1926 Patient was originally treated in the Calcutta School of Tropical Medicine and described by Acton and Napier in the Indian Journal of Medical Research in July, 1927 (Plate) The picture of the patient was subsequently reproduced from this Journal in Manson's Tropical Diseases Ninth edition 1929 The patient was then considered to have resisted all forms of treatment. The treatments adopted by the above observers are not stated

The First Stage of Observation-

This was the stage at which the patient came under the observation of Brahmachari for the first time up to the com mencement of treatment with usea stibamine-the stage of kala azar with dermal lesions due to Leishmania donovani

Patient was suffering from kala azar with fever and enlargement of spleen The peripheral blood gave positive flagellate culture Patient had very well marked nodular eruptions over his face, the trunk and the extremities sides these there were patches of de pigmentation over the body especially the trunk

Blood examination on 28th July 1928 showed -Red blood corpuscles-4 000 000 white blood corpuscles -2 700 haemoglobin-75 per cent Leucocyte count per centages were Polymorphonuclears-61 mononuclears-18 lymphocytes-17 eosinophiles-4 No malarial parasites

Wassermann reaction-negative The nodules in the skin showed Leishmania donovani in smears and flagellates in culture

Patient was put on a course of treatment with urea stibamine from 15th August, 1926

The Second Stage of Observation-

This was the stage during which the patient underwent treatment with urea stibamine and passed toward its end into the stage of dermal leishmanoid with complete sterilization of the internal organs against leishmania

At the beginning of this stage the face and the trunk showed nodular eruptions and patches of de-pigmentation. The eruptions were very similar to those in the first stage. During treatment with urea stibamine the spleen quickly diminished in size, and the fever stopped. The skin eruptions seemed to be resistant to treatment, though they slowly and slightly diminished from the trunk and the extremities. Blood examination on 13th September, 1926, after administration of 1 1g of urea stibamine showed the following —Red blood corpuscles—4,500,000, white blood corpuscles—6,600, haemoglo-bin—75 per cent. Leucocyte count percentages were.—Polymorphonuclears—61, mononuclears—18, lymphocytes—19, eosinophiles—2

Blood culture for flagellates—negative No fever, no enlargement of the spleen at the end of this stage.

At the end of this stage the case was completely cured of kala-azar, but was still suffering from dermal leishmanoid

The Third Stage of Observation-

This was the stage of dermal leishmanoid during which the patient underwent further treatment with urea stibamine and passed toward its end into the fourth stage, when he was completely cured of dermal leishmanoid

We persisted in treatment with urea stibamine and after the further administration of 9 3 g of urea stibamine extending over a period from 13th September, 1926, to 29th August 1927 the patient was completely cured At the end of this stage there was complete sterilization of the tissues against leishmania

The Fourth Stage of Observation-

This was the stage of complete cure and sterilization of all the tissues against leishmania

The patient was examined by Brahmachari on 1st May 1929 about eight months after completion of treatment was then a perfectly healthy man without any enlargement of the spleen and with complete disappearance of all traces of dermal leishmanoid (Plate II)

Plate I represents the appearances of the skin eruptions over the face of the patient as reproduced in Manson's Tro pical Diseases 1929 from Acton and Napier's paper in the Indian Journal of Medical Research July 1927 It is reproduced here for purposes of comparison

Blood examination on 1st August 1927 was as follows -Red blood corpuscles-5 000 000 white blood corpuseles-7 500 haemoglobin-100 per cent Leucocyte count percentages were -Polymorphonuclears-60 0 large mononuclears-3 2 lymphocytes-33 6 eosmophiles-3 2

Culture of the peripheral blood-negative

The case is of great interest as being one in which well marked nodules containing Leishman Donovan bodies were observed in the skin at a stage when the patient was not cured of internal leishmaniasis. The skin eruptions still persistad after the patient had been cured of kala azar and the internal organs had been sterilized against leishmania the case passing into the condition of dermal leishmanoid subsequent history of the case is also of much interest Originally considered to have been absolutely refractory the case was completely cured of dermal leishmanoid after a prolonged course of treatment with urea stibamine Eight

months after the completion of treatment the patient showed no sign whatever of leishmania infection

NB—A word about the nomenclature of the disease Various names have been suggested, one of the latest being " post-kala-azar dermal leishmaniasis" (Acton and Napier. 1927) Very recently, in discussing the various names suggested, the Editor, Indian Medical Gazette (1928), pointed out that the name "post-kala-azar dermal leishmaniasis was a clumsy name, though it defined the condition, the alternative, post-generalised dermal leishmaniasis was certainly worse " The name "leishmanide" suggested by him is not free from difficulties for the following reasons A name ending in ide frequently gives an impression of an amide, such as tryparsamide, and confusion may arise in the mind of the reader as to whether leishmanide is the name of a specific for leishmaniasis or the name of a cutaneous manifestation of infection with leishmania We are therefore inclined to think that on the whole "dermal leishmanoid" is the most appropriate, if not the most convenient, name for the disease At any rate it has now the sanction of usage, and most observers recognise the disease under that name

[N B — This case is the continuation of the one already described in pp 207-209 of this work — Ed]

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STUDIES IN KALA AZAR AND CHEMOTHERAPY OF ANTIMONY

PART II

THE TREATMENT OF KALA AZAR WITH INTRAMUSCULAR INJECTION OF SODIUM N PHENYL GLYCINE AMIDE 4 STIBINATE

In a paper published in the Indian Journal of Medicine June 1926 and in the Calcutta Medical Journal June 1926 Brahmachari and his assistants described a series of eight cases of kala azar successfully treated with intravenous injection of sodium N phenyl glycine amide 4 stibinate (the antimony analogue or tryparsamide)

INTRAVENOUS INJECTIONS

The notes of these published cases are summarized below for reference

Case I—D act 16 was admitted with history of fever for six months. The spleen extended 5 in below the costal arch. Blood culture on NNN medium gave a positive flagellate culture. Patient was treated with intravenous injections of which twenty were given. The fever completely stopped the spleen ceased to be felt below the costal arch, and at the time of discharge blood culture on NNN medium was negative. Increase of weight during treatment—1 st. 1 lb. Dose—0.05 to 0.2 gram.

Result of Blood Examination—

- (1) R B C —1,800,000, W B C —2,500, Hb —35 per cent, on 11th February, 1925, before treatment.
- (2) RBC--4,500,000, WBC--8,400, Hb--85 per cent, on 20th June, 1925, after treatment

Case II — J, æt 10, was admitted on 17th November, 1924, with history of fever for five months. The spleen extended 4½ in below the costal arch. Patient was treated with fifteen intravenous injections. The fever stopped after the third injection. Increase in weight during treatment—1 stone. Dose—0.05 to 0.2 gram

Result of Blood Examination—

- (1) R B C —2,500,000, W B.C —3,500, Hb —45 per cent, on 19th November, 1924, before treatment
- (2) R B C —4,400,000, W B C —9,600, Hb —45 per cent, on 27th March, 1925, after treatment

Case III —S, æt 14, was admitted on 3rd May, 1925, with history of fever for three months. The spleen extended $2\frac{1}{2}$ in , and the liver 1 in below the costal arch. Patient was treated with twelve intravenous injections. The fever stopped after the fourth injection. Dose—0.05 to 0.20 gram

Result of Blood Examination—

- (1) RBC—2,000,000, WBC—1,800, Hb—40 per cent, on 20th June, 1925, before treatment
- (2) R B C —4,200,000, W B C —7,100, Hb —70 per cent, on 12th August, 1925, after treatment

Case IV — J, æt 10, was admitted on 2nd March 1925, with history of fever for four months. The spleen extended to the umbilicus, and the liver was just palpable below the costal arch. He was treated with ten intravenous injections. The temperature came down to normal after the

fourth injecton At the time of discharge the spleen and the liver could not be felt below the costal arch Dosc—0 05 to 0 2 gram

Result of Blood Lxamination-

- (1) R B C —2 500 000 W B C —2 500 Hb —45 per cent on 8th March 1925 before treatment
- (2) R B C -4 000 000 W B C -5 100 Hb -75 per cent, on 10th May 1925 after treatment

Case V —M æt 20 was admitted on 25th May 1925 with history of fever for two years. The spleen extended 6 in below the costal arch. Patient was treated with fifteen intravenous injections. The temperature came down to normal after the fifth injection. At the time of discharge the spleen was felt 2 in below the costal arch. Dose—0.05 to 0.20 gram.

Result of Blood Examination-

- (1) R B C —2 500 000 W B C —2 100 Hb —45 per cent on 27th May 1925 before treatment
- (2) R B C —3 200 000 W B C —6 500 Hb —70 per cent on 17th August 1925 after treatment

Case VI—S are 21 was admitted on 28th February 1925 Both spleen and liver extended 4 in below the costal arch. He was treated with fourteen intravenous injections. The temperature came down to normal after the fifth injection. At the time of discharge the spleen was felt 1 inch. below the costal arch. Dose—0.05 to 0.2 gram.

Result of Blood Examination-

- (1) R B C -2 820 000 W B C 1 800 Hb --30 per cent on 31st March 1925 before treatment
- (2) R B C -3 000 000 W B C -5 000 Hb -65 per cent on 29th May 1925 after treatment

Case VII —B, æt 12, was admitted on 23rd March, 1925. The spleen extended 8 in and his liver 2 in below the costal arch. Patient was jaundiced. He was treated with ten intravenous injections. The temperature came down to normal after the sixth injection. At the time of discharge the spleen could not be felt below the costal arch Increase of weight after treatment—1st. 7lbs. Dose—0.05 to 0.2 gram.

Result of Blood Examination-

- (1) RBC -2,500,000, WBC -2,800, Hb -20 pcr cent, on 26th March, 1925, before treatment
- (2) R B C -4,500,000, W B C -6,200, Hb.-80 per cent, on 10th June, 1925, after treatment

INTRAMUSCULAR INJECTIONS

The above cases showed satisfactory results obtained by intravenous injection in the treatment of kala-azar. As the compound is the antimony analogue of tryparsamide, the possibility of using it intramuscularly with advantage was suggested by Brahmachari. There was for some time considerable difficulty in preparing the compound in a pure state. This was subsequently overcome, and the chances of local irritation after intramuscular injection by the presence of minute impurities eliminated as much as possible

The following are the notes of five cases of kala-azar successfully treated by intramuscular injections. The first case was communicated by Brahmachari to the International Congress of Tropical Medicine and Hygiene, held at Cairo, in December, 1928 and is quoted here for reference

Case I.—Patient, æt 25, came under the treatment of Brahmachari, with history of fever lasting for about six months. The spleen extended 5 in and the lever 3 in below

the costal margin and Leishman Donovan bodies were found on spleen puncture. Blood picture at the commencement of treatment was RBC—3 500 000 WBC—3 500 Hb—45 per cent. Patient was given injections in does of 0.1 to 0.3 gram subcutaneously and intramuscularly. Temperature came down to normal after the fourth injection. After the twelfth injection the spleen could not be felt below the costal margin and no Leishman Donovan bodies could be found on spleen puncture. The blood at the time of communication to the Congress (three weels after the commencement of treatment) showed. RBC—3 500 000 WBC—6 500 Hb—65 per cent. Generally speaking there was very little local reaction at the sents of subcutaneous or intramuscular injection. The patient was completely cured at the time of writing the present paper.

Case II—H C M came under the treatment of Brahmachari on 3rd January 1929 with history of fever of four months duration. Spleen at the time of first observation extended 4 in and liver 2 in below the costal margin. Result of blood examination. R.B. C.—315 000. W.B. C.—3 200. Hb—45 per cent on 10th January 1929. Spleen punch treatment with fifteen intramuscular injections in doses of 0.05 to 0.15 gram twice a week. The temperature came down to normal after the fourth injection. On 7th April 1929 at the time of discharge spleen and liver could not be felt below the costal margin. Blood examination.—R.B. C.—4.500.000. W.B. C.—8.000. Hb.—75 per cent. Peripheral blood culture—negative for flagellates.

The Notes of three cases recently treated in the Chitta ranjan Hospital Calcutta are given below —

Case III —R C J act 10 was transferred from the Cholera Ward to the Kala azar Ward of the Chuttaranjan Hospital on 24th June 1929 His leucocyte count at the

beginning of the treatment was 3,100. The spleen was 4 in below the costal margin. Leishman Donovan bodies were found on spleen puncture. Patient was given altogether eleven injections in doses of 0.05 to 0.1 gram. Temperature came down to normal after the second injection and it continued to be so except for one sharp rise which occurred after the fifth injection, but it rapidly went down to normal. At the time of discharge the leucocyte count was 7,200, and the spleen could not be felt below the costal arch. No Leishman-Donovan bodies could be found on spleen puncture. Increase in weight—1 stone.

Case IV — K D, æt 21, was admitted into the Kalaazar Ward of the Chittaranjan Hospital, with a history of irregular attacks of fever for six months. His spleen was felt 6 in and the liver 2 in below the costal margin. Many Leishman-Donovan bodies were found on spleen pucture Leucocyte count at the beginning of treatment was 3,100. Patient was given thirteen intramuscular injections in doses of 0.1 to 0.15 gram. At the end of treatment spleen and liver could not be felt below the costal margin. Result of blood examination.—R B C —4,600,000, W B C —6,500, Hb —90 per cent. Spleen puncture—negative. Increase in weight—2 stone.

November, 1929, into the Kala-azar Ward of the Chittaranjan Hospital, with history of fever for one month and with spleen enlarged to about $3\frac{1}{2}$ in. and liver 2 in below the costal margin. Spleen puncture showed a large number of Leishman-Donovan bodies. Blood examination at the beginning of treatment.—Leucocyte count—4,500, Differential count polymorphonuclears—50 per cent, lymphocytes—44 per cent, mononuclears—6 per cent, poikilocytes, anisocytes, normoblasts, myelocytes and basophiles present. Patient was put on intramuscular injections in doses of 0.05 to 0.1 gram. Temperature came down to normal after the

fifth injection Patient was given altogether fifteen injec tions in doses of 0.05 to 0.1 g. At the time of writing spleen and liver could not be felt below the costal margin Result of blood examination RBC-4 500 000 WBC-7 500 Hb -80 per cent Spleen puncture-negative

ORSERVATIONS

Sodium N phenyl glycine amide 4 stibinate The antimony analogue of tryparsamide has been successfully used by the intramuscular method in the treatment of kala azar The local irritation caused by it is frequently slight same drug has also been successfully used by the intravenous method in the treatment of kala azor

The maximum tolerated dose in the case of the white rat given intravenously is 0.3 gram per kg of body weight

Many antimony preparations have been used from time to time intramuscularly for the treatment of kala azar A Reference to their use is given below -

- Castellani s solution of tartar emetic 1
- 2 Brahmachari s hyperacid antimonyl tartrate
- 3 Sodium antimonyl tartrate
- Acetyl p amino phenyl stibinate of sodium (Stibe 4 nyl)
 - 5 Urea stibamine
 - Neo stihosan

Many of the above compounds have been found to give rise to much local irritation and up to now none has yet been popular for the purpose of intramuscular injection

In view of the fact that the present compound is allied to tryparsamide its value in the treatment of kala azar by the intramuscular route cannot be over estimated lts use is indicated in individuals with thin veins or in whom intra

venous injections of any antimony compound are followed by severe constitutional symptoms The compound was first discovered by Brahmachari in 1922

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STUDIES IN KALA AZAR AND CHEMO THERAPY OF ANTIMONY

PART III

OBSERVATIONS ON ANTIMONY IN THE SPLEEN CELLS
OF ANIMALS INFECTED WITH LEISHMANIA DONOVANI

The pathology of the spleen in kala azar is an important subject for research. In experimentally infected animals the spleen may show little or no fibrous tissue development but contains in heavily infected cases a very large number of cells harbouring Leishmania Donovani. These cells possess the property of picking up particles of antimony when it is introduced into the circulation in a state of fine subdivision as will be seen from the experiments described below.

Metallic antimony in a state of fine subdivision was prepared according to the method of Plimmer (1911) in the following way

By dissolving antimony trichloride or other salts of antimony in hydrochloric acid or other acids diluted with water or an aqueous solution of an organic acid such as tartanc acid and adding zinc to the solution the antimony is precipitated in a finely divided state. The proportions employed are 50 grams of antimony trichloride 100 c cm of concentrated hydrochloric acid diluted with 200 c cm of water or 200 c cm of 2 to 5 per cent tartanc acid solution and 25 to 30 grams of zinc. When the zinc is completely dissolved the antimony is filtered off and

washed with an aqueous solution of an organic acid, such as tartaric acid, until free from chlorides, and then with water till free from acid. The antimony thus obtained is pure and in a finely divided state."

One per cent suspension of metallic antimony prepared in the above way was made in normal saline in a test-tube and the heavier particles were allowed to settle for ten to fifteen minutes at the bottom of the tube

The supernatant suspension was injected intravenously into healthy as well as leishmania-infected mice in doses of 20 c cm. per kilo of body weight

The animals were killed with coal gas at intervals of two hours, two days and seven days after injection, and sections of the organs were then prepared

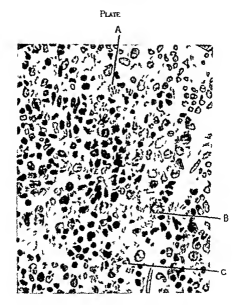
No antimony was found inside the cells of the spleen two hours after injection. On the other hand, two days and seven days after injection antimony was found inside certain cells of the lungs and spleen

The section of spleen shown here (see Plate) is that of a leishmania-infected mouse killed forty-eight hours after intravenous injection of metallic antimony in doses of 20 c cm per kilo of body weight on two successive days, and stained with iron-hæmatoxylin. On careful examination the following types of cells were found inside the spleen of the mouse. (Some of the cells are shown in the Plate.)

- 1 Large cells with faintly stained cytoplasm containing leishmania as well as antimony particles
- 2 Cells containing antimony in a diffuse (probably colloidal) state, which may or may not harbour leishmania
- 3 Cells of moderate size containing a large number of leishmania as well as particles of antimony
- 4 Certain small cells containing particles of antimony, although they do not contain any leishmania
- 5 Large cells with faintly stained cytoplasm containing very few or no antimony particles, but full of leishmania

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- 6 Certain fairly large cells with particles of antimony and degenerate leishmania inside them
- 7 Giant cells containing very few or no antimony particles or leishmania
- 8 Cells with fairly large and well stained nuclei con taining neither antimony nor leishmania

In certain places there were particles of antimony free in blood spaces inside the spleen. We have further observed that cells harbouring the largest number of leishma nia do not necessarily pick up the largest number of particles of antimony

Some of the above cells are probably of the nature of clasmatocytes

From the above it will be seen that antimony was present inside the cells of the spleen in two forms (1) as a diffuse brownish yellow stain and (2) as black granules. We consider that the diffuse staining is due to the metallic antimony being converted into a colloidal form by the cells of the spleen in the process of its internal metabolism before it is converted into a soluble compound

These observations appear to have an important bearing upon the action of antimony in the treatment of kala azar

Levaditi has propounded a general law with reference to all the members of the nitrogen family of elements occupying Group V of Mendeleff's periodic table—arsenic antimony variadium bismuth etc. They or their compounds exhibit paraciticidal properties after they have been acted upon by certain cells. It is probable that some of these cells are clasmatocyte cells giving rise in the case of bismuth to a toxalbumin the bismoxyl of Levaditi which possesses destructive power against the Treponema pallidum

It has been suggested by Brahmacharı elsewhere that an antimony compound in order that it may be of thera peutic value must be converted in the tissues into a

compound containing the radical -Sb=0 in the reactive stage Chemically, some of the therapeutic antimony compounds contain the radical -Sb=0, and the therapeutic bismuth compounds the radical $-B_1=0$. It is likely that bismoxyl contains the radical $-B_1 = 0$ in the reactive stage, and that a corresponding antimony compound which has been called stiboxyl (Brahmachari, 1928) is probably formed in the case of antimony. It has been observed that metallic bismuth, finely subdivided, is more suitable for the production of bismoxyl when administered intramuscularly than in the form of a chemical compound. The same could be expected of metallic antimony, but for the fact that when injected intramuscularly it gives rise to so very severe local irritation that it is unsuitable for inframuscular injection for therapeutic purposes On the other hand, metallic antimony injected intravenously is one of the most powerful antimonials in the treatment of kala-azar, as was shown by Brahmachari (1915) some years ago It has been observed by Meleney that in kala-azar clasmatocyte tissue is developed as a Probably this reticulo-endothelial system tissue reaction gives rise to the production of stiboxyl in the spleen and If this view be correct, then it may be conelsewhere cluded that individual cases will get beneficial results from the use of antimony compounds proportional to the reaction of the reticulo-endothelial system. The following two requirements are therefore necessary for an antimonial to be effective, namely (1) the development of the clasmatocytes, and (2) the introduction of an antimony compound with which they can combine for the development of stiboxyl and the ability of the clasmatocytes to metabolise the antimon'al and give rise to one which will act upon the Herein lies the value of the different antimonials in the treatment of kala-azar This will explain why with the same antimony compound, one individual is cured much more quickly than another after its administration It is the

development of these tissue cells and their ability to meta bolise the compound used that should be aimed at in the treatment of the disease. It is clear from the above obser vations that after intravenous administration of metallic antimony into an animal experimentally infected with kala azar the leishmania come into closest contact with the particles of antimony inside certain tissue cells and are subsequently destroyed by a highly reactive antimony compound formed inside them in the process of metabolism inside these cells How these cells may be developed and how their ability to metabolise antimony compound picked up by them may be increased is a subject for further research

OBSERVATIONS

Metallic antimony injected intravenously in a state of fine subdivision into leishmanin infected mice is picked up inside the spleen by cells that harbour leishmania and an antimony compound is developed in ide the cells during the process of metabolism which kill the leishmania certain cells inside the spleen metallic antimony is found in a diffuse state and this is probably the stage in which the solid finely divided antimony is converted into colloidal particles before passing into complete solution harbouring the largest number of leishmania do necessarily pick up the largest number of particles of antimony Degenerate leishmania have been observed inside cells containing particles of antimony

The senior author's grateful thanks are due to Major Shortt Director Kala azar Commission Assam for providing him with leishmania infected mice. Without his aid it would not have been possible to conduct the above work. He is also grateful to Dr Baini Prashad of the Zoological Survey of India for the beautiful photo micrograph of the section of

spleen of the infected mouse which is attached to the present paper

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Daniellson

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STUDIES IN KALA AZAR AND CHEMO THERAPY OF ANTIMONY

PART IV

FURTHER OBSERVATIONS ON THE ANTIMONY LADEN CELLS OF SPLLEN AFIER INTRAVENOUS INJECTION OF METALLIC ANTIMONY IN A STATE OF FINE SUSPENSION IN EXPERIMENTAL ANIMALS

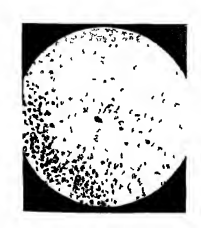
In our last paper communicated to these Transactions (Vol XXIII No 5) we found that when metallic antimony in a fine state of subdivision is injected intraven ously into a leishmania infected mouse many of the antimony particles are picked up inside the spleen by cells containing leishmania and others which do not contain them. These cells are probably clasmatocytes

In the present paper the results of observations on healthy mice after intravenous injections of finely divided metallic antimony are recorded. The same procedure as was adopted in our former paper was followed in our present observations. One per cent suspension of finely divided metallic antimony prepared according to the method of Plimmer was made in normal saline solution and then allowed to settle for ten to fifteen minutes and the superna tant suspension was injected into the dorsal vein of the tail of a mouse in doses of 20 c cm per kilo. The injection was repeated after twenty four hours and the animal killed forty eight hours after the first injection.

In sections of the spleen of these experimental animals we find that the particles of antimony are picked up by cells

similar to those in the case of leishmania-infected mice, with the difference that in the case of healthy mice a larger number of particles of antimony are extracellular forty-eight hours after the first and twenty-four hours after the second injection than in the case of infected mice. This is explained by the fact that in the infected mice, a larger number of cells of the nature of clasmatocytes, which are capable of taking up leishmania as well as antimony particles, are developed in them as a result of tissue reaction than those present in healthy mice Further, it has been observed that in the case of infected mice these cells are much larger in size, and the particles of antimony inside them are more dispersed than in the case of healthy mice, in which they are present in a more compact state (Compare the Photomicrograph in our previous paper on the subject with that in the present paper) This may be explained by assuming that the leishmania-laden clasmatocytes differ in the degree of their functional activity from those present in healthy mice

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STUDIES IN KALA AZAR AND CHEMO THERAPY OF ANTIMONY

PART V

THE TREATMENT OF RESISTANT CASES OF DERMAL LEISHMANOID*

In treating cases of kala azar with antimonial pre parations the authors have met with the following types (1) Those which quickly yield to antimonial treatment (2) those which resist for a considerable period and slowly yield to treatment (3) those which seem to be extremely or abso lutely resistant (4) those which relapse after insufficient or improper treatment and are either very resistant to subse quent treatment or quickly yield to it. Thus patients vary in their response to treatment while the possibility of antimony resistant leishmania which may be present from the very b ginning or develop during treatment has to be con sidered Relapses occur after too early abandonment of treatment and also but rarely in cases which from the marked improvement in the general condition and in the blood picture and freedom from fever for some months appear to have been cured

As regards dermal leishmanoid contrary to the opinion of Muir (1930) it is found that most cases take much

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longer to cure than cases of kala-azar This fact is significant, for the disease is more common than was supposed when it was first discovered (1922) The unsightly nature of the lesions makes the patients very unhappy as well as objectionable to others. Furthermore, such cases are probable sources of infection, so that the need for a rapid cure cannot be over-emphasized. A most resistant case of dermal leishmanoid which was originally considered by Acton and Napier as absolutely refractory to treatment, has been previously recorded in this Journal (1929) The course of treatment adopted was not indicated by these authors In a more recent communication, however, reference is made to the case by Napier and Haldar (1930) It is stated that between fifty and sixty injections of different antimony compounds were given Urea stibamine alone is named, and of this the patient is stated to have had twelve injections As noted (1929), this case was subsequently cured by a very prolonged course of urea stibamine It is evidently important to discover methods for reducing the course of treatment.

This paper refers to those cases of dermal leishmanoid which show little or no improvement after continuous antimonial treatment extending over a period of three months or more. The type of case recently mentioned by Napier as having been cured after the usual course for the treatment of kala-azar is not dealt with here, though such has also been observed by us

On the assumption that skin lesions would respond most quickly if antimony were introduced through the skin, Brahmachari conceived the idea of combining ointment of metallic antimony with intravenous injection of urea stibamine. Six cases have now been treated in this way, and the results appear so far to be much more satisfactory than treatment by the intravenous route alone. The notes of one of the most resistant of these cases are given here

The patient came under observation in January 1929, with well marked dermal lesions due to Leishmania donovani He gave a history of having had kala azar about two years previously and of being cured after antimonial treatment After forty injections of urea stibamine in doses of 0 I to 0 82 gram he was much improved Treatment was discontinued for six months after which there were well marked nodules on his face and marked patches of de pigmentation all over the body. The patient then received during two months three injections of urea stibamine and twelve of neo stibosan but as improvement was very slight a combined treatment of metallic antimony inunction (5 per cent) and intravenous injection of urea stibamine was insti tuted After twenty seven injections had been given all the nodules on the skin had disappeared. The patient is still under our observation

In addition to the above treatment we have in a few cases tried berberine sulphate—a drug used by old writers for the treatment of enlarged spleen in India O Shaugh nessy recommended it for the treatment of ague and remittent fevers while in the early days of the antimony treatment of kala azar Brahmachari used this drug intravenously in the treatment of kala azar without any benefit (1916) Berberine has however had a long standing reputation for being useful in the treatment of oriental sore and is obtainable in the market in a crude state under the popular name of rasaut Recently acid berberine sulphate has been used with success in the treatment of the same disease both as an ointment and by infiltration of the sore It is now obtainable under the name of orisol We have used this in a few cases of dermal leishmanoid for the treatment of the de pigmented patches but up to the present no benefit has been obtained from the use of this drug intra venously or subcutaneously

Metallic antimony used by us was in the form of an

impalpable powder prepared by the method of Plimmer, which has already been described by us in this Journal (1930). The ointment used contains 5 per cent of metallic antimony in equal parts of lanoline and vaseline. It is gently rubbed over the affected parts of the skin for ten to fifteen minutes daily till the skin lesions disappear. In two cases, we injected 0 125 to 0 25 c cm of a 5 per cent solution of urea stibamine into some of the nodules, and this was followed in a few days by marked shrinkage without any local irritation.

Conclusions

In order to shorten treatment, cases of dermal leishmanoid should be treated by metallic antimony inunction combined with intravenous injection of a therapeutic aromatic pentavalent antimonial, which, in our cases, consisted mostly of urea stibamine

References

STUDIES IN KALA-AZAR AND CHEMO THERAPY OF ANTIMONY

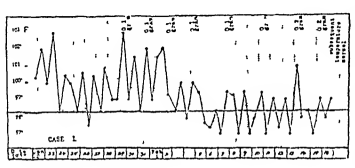
PART VI

TREATMENT OF KALA AZAR WITH INTRAMUSCULAR INJECTION OF SODIUM SULPHOMETHYL STIBANILATE

In these Transactions Brahmachari and his co workers de scribed in 1930 a series of cases of kala azar successfully treat ed with intramuscular injection of sodium N phenyl glycine amide 4 stibinate the antimony analogue of tryparsamide Excepting this arsenic compound the organic therapeutic arsenicals are generally derivatives of arsenobenzene the other hand the organic therapeutic antimonials are gene rally derivatives of p stibanilic acid Among the therapeutic aromatic arsenicals sodium methylene sulphonic acid deri vative of dioxy diaming arsenobenzene or di sodium dioxy diamino arsenobenzene methylene sulphonate has been suc cessfully used intramuscularly in the treatment of syphilis under the various trade names of thio sarmine sulfarsenol and it occurred to the writer to give a trial to sodium methylene sulphonic acid derivative of stibanilic acid or sodium sulphomethyl stibanilate in the treatment of kala azar As stibinobenzene compounds are unstable they have not yet come into use in therapeutics and no attempt was made to synthetize them

The synthesis of sodium-sulphomethyl-stibanilate has been successfully effected in the Department of Chemistry of the Brahmachari Research Institute, Calcutta—It is a greyish white heavy powder, freely soluble in water and its solution is very faintly acidic to litmus—Its antimony content—is 24.5 per cent

The following is a series of cases of kala-azar successfully treated with intramuscular injection of this compound.



Case I—A K
M, male, æt 12,
was admitted into Brahmachari's
ward in the Carmichael Medical
College Hospitals with history
offever for six
months The

spleen was hard and extended 5 in. and liver extended $1\frac{1}{2}$ in below the costal arch—Leishman-Donovan bodies were found on spleen puncture—Patient was given—21 injections of the compound intramuscularly in doses of 0.1 to 0.3 gramme, at first every other day and subsequently twice a week—Effect of treatment on temperature is shown in the accompanying Chart—At the time of discharge, spleen and liver could just be felt below the costal arch

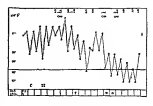
Total dose-4 6 grammes

Result of Blood Examination—

Before treatment R.B C \longrightarrow 3,375,000, W B C \longrightarrow 2,500, Hb. \longrightarrow 45 per cent

After treatment R B C -4,100,000, W.B C -6,380, Hb.-80 per cent

Case II -A C M male aet 12 was admitted into Brahma charts ward in Carmichael Medical College Hospitals with history of feve for eight months The spleen was hard and extended 2 in and



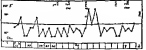
liver also extended 2 in below the costal margin. Leishman Donovan bodies were found on spleen puncture Patient was given fifteen injections of the compound intramuscularly in doses of 0.05 to 0.2 gramme generally every day and sometimes every other day Effect of treatment on tem perature is shown in the accompanying Chart At the time of discharge spleen and liver could not be felt below the costal arch

Total dose-2 11 grammes

Result of Blood Examination-

Before treatment R B C -2 240 000 W B C -2 300 Hb --45 per cent

R B C -3 500 000 W B C -6 250 After treatment Hb -60 per cent



Case III -S female aet 26 was admitted into the ward of Dr P Nandi Physician Car michael Medical College Hospitals with history

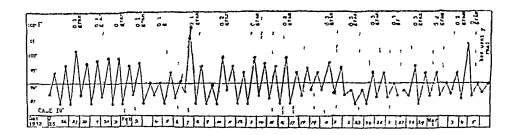
of fever for one year and by his kind courtesy the patient was put under my treatment. The spleen was hard and extended 4½ in and liver extended 1½ in below the costal margin. Leishman-Donovan bodies were found on spleen puncture. Patient was given thirteen injections of the compound intramuscularly in doses of 0.1 to 0.3 gramme generally every other day. Effect of treatment on temperature is shown in the accompanying Chart. At the time of discharge, spleen and liver could just be felt below the costal arch.

Total dose-3 7 grammes

Result of Blood Examination-

Before treatment R B C -2,290,000 , W B C -3,450 After treatment R B C -4,000,000 . W B C -5,624 , Hb -55 per cent

Case IV—MB, male, set 30, was admitted into the Carmichael Medical College Hospitals, with history of fever for about a year. There was bleeding from the gums and patient had frequent attacks of severe epistaxis. There was presence of copious albumin in the urine. There was sedema in the extremities. Spleen was hard and extended



4 in and liver extended 3 in below the costal margin Leishman-Donovon bodies were found on spleen puncture Patient was given seventeen injections of the compound intramuscularly in doses of 0.1 to 0.3 gramme generally every other day Effect of treatment on temperature is shown in the accompanying Chart. At the time of discharge spleen and liver could just be felt below the costal arch and there was no albumin in the urine and no Leishman Donovan bodies were found on spleen puncture

Total dose-3 75 grammes

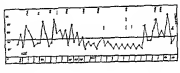
Result of Blood Examination-

Before treatment R B C -- 2 300 000 W B C -- 2 300 Hb -- 40 per cent

After treatment R B C —3 000 000 W B C —5 000 Hb —50 per cent

Case V -A C

M æt 36, was admitted i n t o the Carmichael Medical College Hospitals with history of fever



for about a year

The spleen was hard and extended 23 in and liver extended 1 in below the costal margin. Patient was given nine injections of the compound intramuscularly in doses of 0 1 to 0.5 gramme at first every other day and subsequently twice a week. Effect of treatment on temperature is shown in the accompanying Chart. At the time of discharge spleen and liver could just be felt below the costal arch.

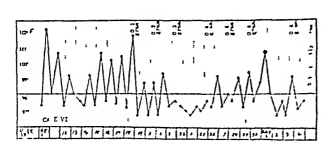
Total dose-3 6 grammes

Result of Blood Examination-

Before treatment RBC-3000000 WBC-2812 Hb-55 per cent

After treatment R B C -- 3 250 000 W B C -- 6 250 Hb -- 83 per cent

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Case VI — KD, male, æt 26, was admitted into the Carmichael Medical College Hospitats, with history of fever for two years The spleen

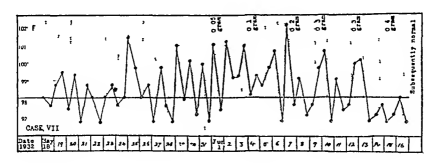
was hard and extended 5 in and liver extended 2½ in below the costal margin. Leishman-Donovan bodies were found on spleen puncture, and in addition there were crescents in the blood Patient was given eleven injections of the compound intramuscularly in doses of 0°1 to 0°4 gramme, at first every other day and subsequently twice a week, together with a course of treatment with quinine Effect of treatment on temperature is shown in the accompanying Chart At the time of discharge, spleen and liver could just be felt below the costal arch, and no Leishman-Donovan bodies were found on spleen puncture

Total dose—3 7 grammes

Result of Blood Examination—

Before treatment R B.C -2,850,000 W B C -2,720 After treatment R B C -3,250,000 WB C -6,250, Hb -60 per cent

Case VII —R M-, male, æt 20, was admitted into the Čarmichael Medical College Hospitals, with history of fever for six months The spleen was hard and extended



7 in and liver extended 1 in below the costal margin Leishman Donovan bodies were found on spleen puncture Patient was given eleven injections of the compound intramuscularly in dose of 0.05 to 0.4 gramme generally twice a week. Effect of treatment on temperature is shown in the accompanying Chart. At the time of discharge spleen and liver could not be felt below the costal arch.

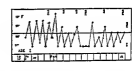
Total dose—3 4 grammes

Result of Blood Examination-

Before treatment RBC-1 450 000 WBC-2 188 Hb-45 per cent

After treatment R B C -3~000~000 W B C -5~000 Hb -60~per~cent

Case VIII—N male act 32 was admitted into the Carmichael Medical College Hospitals with history of fever for six months The spleen was hard and extended 3½ in



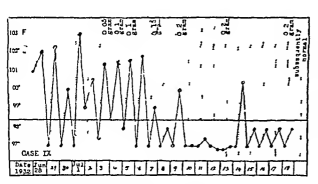
and the liver extended 1½ in below the costal margin. Leish man Donovan bodies were found on spleen puncture. Patient was given thirteen injections of the compound intramus cularly in doses of 0.05 to 0.3 gramme at first every other day and subsequently twice a week. Effect of treatment on temperature is shown in the accompanying Chart. At the time of discharge, spleen and liver could not be felt below the costal arch.

Total dose-2 5 grammes

Result of Blood Examination-

Before treatment RBC -2750000 WBC -2500 Hb -50 per cent

After treatment RBC-3 100 000 WBC-6 688 Hb-65 per cent

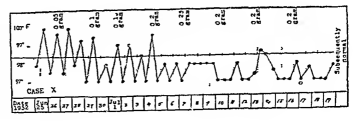


Case IX.—N.CP, male, æt 30, was admitted into the Carmichael Medical College Hospitals, with history of fever for one year He had attacks of diarrhæa

and cough There was presence of albumin in the urine. The Spleen was soft and tender extending 1½ in below the costal margin. Liver also extended 1½ in below the costal margin. Leishman-Donovan bodies were found on spleen puncture. Patient was given seven injections of the compound intramuscularly in doses of 0.05 to 0.2 gramme, at first every day and subsequently every other day and twice a week. Effect of treatment on temperature is shown in the accompanying Chart. At the time of discharge, spleen and liver could not be felt below the costal arch, and there was no albumin in the urine.

Total dose-

Case X—P
PS, male, æt
26, was admitted into the surgical wards
of the Carmichael Medical
College Hospi-



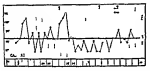
tals for treatment of right perinephritic iliac abscess and subsequently transferred to Brahmachari's ward for treatment of kala-azar. The spleen was hard and extended $3\frac{1}{2}$ in and liver extended $1\frac{1}{2}$ in below the costal margin. Leishman-Donovan bodies were found on spleen puncture. Patient was given eight injections of the compound intramuscularly in doses of 0.05 to 0.2 gramme generally twice a week. Effect of treatment on temperature is shown in the accompanying Chart. At the time of discharge spleen and liver could not be felt below the costal arch.

Total dosc-1 35 grammes

Result of Blood Examination-

Before treatment R B C -2 400,000 W B C -3 500 Hb -50 per cent

After treatment R B C -- 3 800 000 W B C -- 8 750 Hb -- 65 per cent



Case \(\formula\) - B \(\hat{N}\) C male act 35 was ad mitted into Brahma charis ward in the Chittaranjan Hospital with history of fever

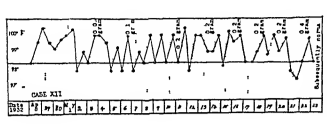
for three months The spleen was hard and extended 41 in and liver extended 2 in below the costal margin Leishman Donovan bodies were found on spleen puncture Patient was given eight injections of the compound intransuscularly in doses of 0 I to 0 2 gramme, at first twice a week and subsequently every other day Effect of treatment on temperature is shown in the accompanying Chart At the time of discharge spleen and liver could not be felt below the costal arch

Total dosc-1 4 grammes

Result of Blood Examination—

Before treatment RBC-2 200 000 WBC-1 550 Hb-45 per cent

After treatment R B C -- 30 00 000 W B C -- 7 800 Hb -- 55 per cent



Case XII—A M, male, æt 26, was admitted into the Chittaranjan Hospital, with history of fever for six months

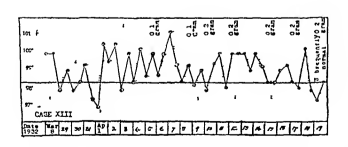
spleen was hard and extended 4 in. below costal margin Liver was not palpable. Leishman-Donovan bodies were found on spleen puncture. Patient was given nine injections of the compound intramuscularly in doses of 0.05 to 0.4 gramme, at first twice a week and subsequently every other day. Effect of treatment on temperature is shown in the accompanying Chart. At the time of discharge, spleen could not be felt below the costal arch.

Total dose—1 95 grammes

Result of Blood Examination—

Before treatment RBC.-1,300,000, WBC-1,800, Hb-35 per cent

After treatment RBC-1,700,000, WBC-6,000, Hb-45 per cent



MNM, male, at 20, was admitted into the Chittaranjan Hospital, with history of fever for some

months The spleen was hard and extended 4 in below the costal margin Liver was not palpable Leishman-Donovan bodies were found on spleen puncture Patient was given fifteen injections of the compound intramuscularly in doses of 0 Lto 0 4 gramme generally every other day and sometimes twice a week. Effect of treatment on temperature is shown in the accompanying Chart. At the time of discharge spleen could just be felt below the costal arch.

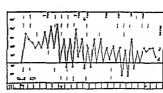
Total dose = 2 85 grammes

Result of Blood Examination—

Before treatment RBC—1 700 000 WBC—1 550 Hb—35 per cent

After treatment R B C -2 000 000 W B C -5 700 Hb -50 per cent

Case XII —
U male set 35
was admitted into
the Chittaranjan
Hospital with his
tory of fever for
six months The
spleen was hard



and extended 6 in and the liver extended 2 in below the costal margin. Leishman Donovan bodies were found on spleen puncture. Patient was given nine injections of the compound intramuscularly in doses of 0 I to 0 4 gramme at first twice a week and subsequently every other day. Effect of treatment on temperature is shown in the accompanying Chart. After completion of treatment spleen and liver could not be felt below the costal arch. Patient is now under observation in the hospital.

Total dose-1 8 grammes

Result of Blood Examination-

Before treatment R B C -- 2 300 000 W B C -- 1 550 Hb -- 50 per cent

After treatment R B C -3,000,000; W B C -5,744, Hb -65 per cent

Case XV—KPM, male, æt. 15, was admitted into the Carmichael Medical College Hospitals, with history of fever for seven months. The spleen was hard and extended 7 in and liver extended 4 in below the costal margin Leishman-Donovan bodies were found on spleen puncture Patient was given ten injections of the compound intramuscularly in doses of 0.05 to 0.3 gramme twice a week At the time of writing this paper, the spleen extended 3 in below the costal margin and the patient was still under treatment

Temperature—Apyrexial throughout the course of treatment.

Total dose—1 9 grammes

Result of Blood Examination-

Before treatment R B C -2,500,000, W B C -2,500, Hb -40 per cent

At the time of writing RBC-2,520,000, WBC-3,500, Hb-60 per cent

OBSERVATIONS

The toxicity of sodium-sulphomethyl-stibanilate is low lts maximum tolerated dose is 0.4 per kilo of body weight in the case of white rats given intravenously. It has been successfully used in the treatment of kala-azar by intramuscular injection. Generally speaking no local or constitutional symptoms have been observed after its use. It has been injected up to a dose of 0.4 gramme. One injection of 0.5 gramme was given to one patient without any untoward results. In one case complicated with nephritis, ædema, and epistaxis, no untoward results followed its use. Originally, the compound was used in doses of 0.1 to 0.2 gramme intramuscularly, but it has been observed that the dosage

can be increased in an adult from 0.2 to 0.4 gramme without any constitutional symptoms. It has been used every other day in some cases and twice a week in others.

Other antimony compounds previously used inframuscularly in the treatment of kala azar have been dealt with in an earlier paper by the author and coworkers and therefore need not be mentioned here.

FORMS OF PYREXIA DUE TO LEISHMAN-DONOVAN BODIES

(1) High intermittent pyrexia —

Fever may or may not be attended with rigors (Chart No 1)

(2) Irregular intermittent pyrexia —

This type of fever is commonly seen. The fever may or may not be attended with rigors.

(Chart No. II.)

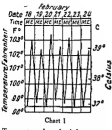
(3) Double quotidian pyrexia with double intermission during 24 hours —

There is rise of temperature towards very early morning followed by intermission before noon. There is a second rise in the evening followed by intermission before midnight (Chart No. III.)

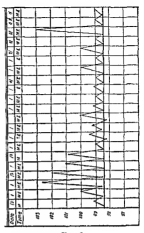
(4) Double quotidian pyrexia with single intermission during 24 hours —

There is a rise of temperature towards very early morning followed by intermission as in the above. There is a second rise in the evening which is followed by remission and not intermission before midnight. (Chart No. IV.)

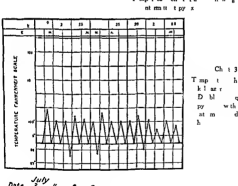
- (5) Intermittent pyrexia with irregular periods of apyrexia
 - (6) Remittent pyrexia



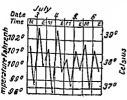
Tmprtr hart f kala azər c sh w ng h gh int m tt nt pyrex



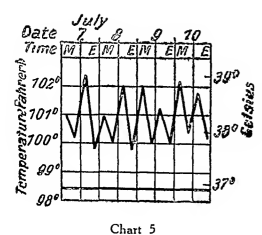
Chat 2 Tmprtu ch t fa gul hwg



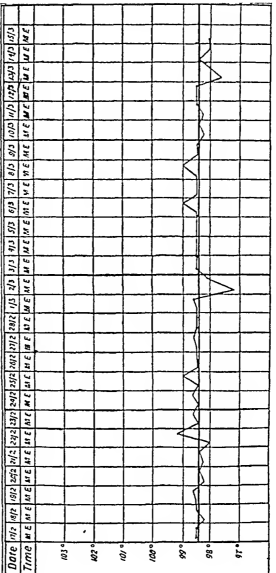
hwng gu td n ld b diw d ng 24



kal azarca Doubl qu tid an py ex: with sing! ant mm ss n an 24 h urs.



Temperature chart of a kala azar case showing Double remittent pyrenia



Temperature chart of a kala-az-1r case showing an almost apyretic temperature

Chart 7

Temperature chart of a kala-azar case showing combined intermittent and remittent pyrexia resembling hectic

(7) Double remittent pyrexia -

There is a rise of temperature towards very early morning followed by remission before noon. There is a second rise in the evening followed again by remission before midnight. The temperature does not come down to normal (Chart No. V.)

(8) Combined intermittent and remittent pyrexia resembling heetic —

Patients may have this type of temperature for a long time (Chart No VI)

(9) The presence of Leishman Donovan bodies is not necessarily associated with much pyrexia (Chart No VII) There may be very slight rise of temperature for some time *

The great peculianty of the pyreixa due to the Leish man Donovan bodies is its variable nature. The various types may be combined in one and the same patient. There was a case who had at first low intermittent fever for some time then high intermittent fever and then remittent fever. No explanation has as yet been offered of this variability of the temperature curve. There may also be variable periods of apyrexia in the course of the disease though the parasites may still be present in the spleen. The double remittent type of pyrexia may pass into the simple remittent type.

Tplqtd py h 1 b td m k l d m tm tl py m y mbl that f pyæm -Ed]

A CONTRIBUTION TO THE STUDY OF FEVERS DUE TO LEISHMAN-DONOVAN BODIES

The discovery of the Leishman-Donovan bodies in the blood of the spleen of patients suffering from what used to be called "malarial cachexia" has revolutionized our ideas about the causation of this disease Until very recently, all cases of enlarged spleen with anæmia, wasting and a history of fever resembling that of true malaria and coming from more or less malarious places, used to be called "malarial cachexia "Six years ago, while studying the malarial fevers in the Calcutta Medical College Hospitals, I used to find a number of cases with enlarged spleen, in whose blood no malarial parasites were found Still one used to call such cases malarial, because their resemblance to malaria was great, though at the same time differing from it in not being amenable to quinine or only slightly so It is, I think, this confusion of malarial fevers with those due to the Leishman-Donovan bodies, that has led many physicians in Bengal to believe that there are two kinds of malarial fever, one amenable to guinine and the other not Cases of true malarial fever not yielding to quinine given in sufficient doses and for a sufficient length of time must be rare. Of course there are cases in which quinine may not be absorbed from the gastro-intestinal tract, but such cases are far from being common One such case I shall cite to you A girl, about 7 years old, was treated by me at Dacca, for intermittent

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^{*} Read at the Calcutta Medical Club, 6th September, 1906

fever coming on with ague every day with slightly enlarged spleen and a tender liver. Her blood showed the presence of malignant tertain parasites. She was given large doses of guinine but there was no disappearance of the fever nor were there any symptoms of quininism quinine was given in solution still I believed that it was not absorbed into the system and my suspicion was corroborated by my being able to detect its presence in the fæces fact is not strange because we know that in another disease viz cholera medicines are not absorbed and may remain inside the stomach unabsorbed for days together and then show suddenly poisonous symptoms in the stage of re action Leaving however these exceptional cases in which quinine is not absorbed into the system we may say almost with certainty that one of the surest tests for malarial fever is its amenability to quinine This fact has been amply proved by the discovery of the Leishman Donovan parasites which are responsible for a large number of cases of fever simulating malaria These latter have resisted quinine giving it the bad reputation of not being infallible in the treatment of malaria

When I first began to study cases of malanal cachexia in the Calcutta Medical College Hospitals it struck me that such cases were much more common and had a much worse prognosis in India than in America I used to explain to myself their more frequent occurrence being due to less frequent use of quimne in India But when I found that they never showed any malanal parasites in the blood that their fever could not be checked by quinine that a very large number of them ended fatally and many of them had cancrum oris and dropsy it struck me that their cause was something that was not known to us

The fever due to Leishman Donovan bodies has been termed cachexial fever or cachectic fever by Major Leonard Rogers But as the disease may exist for a variable period without any fever 1 thml. it may be more appropriately

called leishman-donovan disease or tropical cachesial disease, a term though cumbrous is perhaps more accurate

It has not as yet been worked out how the parasites enter the system. The disease is frequently found in places where true malarial fevers occur. It is probably propagated through water or through the mosquito or the parasite may enter the system through the gastro-intestinal tract.

As regards the carly symptoms of the disease our knowledge is still very limited. The cases that we meet with in the hospital wards are those that have suffered for a long time. I give here a summary of the early symptoms of the disease as gathered from the patients

In one class of cases we get a history of intermittant attacks of fever, quotidian for some time and later on becoming irregular with or without rigors, lasting for several months and not benefited or only temporarily benefited by quinine. In a second class there is no history of any previous attacks of fever but the spleen becomes slowly and steadily enlarged. In a third class there is a history of attacks of low fever, continued for several months with a progressive enlargement of the spleen and not benefited by any treatment. In a fourth class there is history of one or two attacks of remittent fever more or less resembling typhoid. In a fifth class the patient gives history of gastrointestinal troubles with dysenteric or diarrhœic attacks followed by cedema of the lower extremities and attended from time to time with ague-like attacks of fever.

In all these cases, the patient invariably says that there was very slight or no benefit from quinine

One of the most acute and fatal cases that I have ever seen was a child whom I treated some time ago. She was about 4 years old, belonged to a rich family and was brought back to Calcutta from Eastern Bengal where she had been taken to from Calcutta and kept about a month in a malarious district. She was suffering with high remittent

fever lasting for about a week. She had cancrum oris on the 4th day of the fever. When I examined her I found her anæmic with a large portion of the left cheek gan grenous and with slight enlargement of the spleen and slight bronchitis. The fever was of a remittent type. The cancrum oris spread very quickly and the patient died five days after being brought to Calcutta. I might remark here that I had seen the child shortly before her leaving Calcutta. in a per feetly healthy condition.

I have observed eight different types of pyrexia in this disease. These were published in the Indian Medical Ga ette. January 1906. [They will not be reproduced here—Ed.]

The great peculiarity of pyrexia in this disease is its variable nature. As shown before the various types may be combined in one and the same patient. No explanation has as yet been offered of this variability of the temperature curve. Due to some unknown cause the temperature of the patient may remain perfectly normal for several days or weeks though the parasites may be seen in large numbers in the spleen.

A very common symptom in this disease is cedema of the lower extremities—It is sometimes an early symptom of the disease—On the other hand it may be absent even up to the last—It is not easy to explain the cause of the cedema. The cedema is not due simply to aniemia because sometimes it is a very early symptom of the disease and then we would notice it more markedly in true malarial cachexia which is attended as Major Rogers has pointed out with more aniemia.

I show you here the pl-otographs of two cases taken a few days before death one of whom had a very well marked ædema and the other almost none at all Both were advanced cases being anæmic having no albumen In the urine and having no abnormality in the heart It has struck me that we can differentiate the disease into two classes, one in which cedema is a marked symptom and the order in which it is least marked. This would remind one of the two types of beriberi that have been observed, the wet and the dry varieties. Cases in which cedema of the lower extremities is completely absent is more rare than those in which it is present.

Gastro-intestinal symptoms are also often noticed in this disease. Attacks of dysentery or diarrhoea are often present and sometimes there may be a most intractable attack of dysentery which may end fatally.

Wasting and anæmia are marked symptoms of the disease. There is a peculiar facies which enables one to suspect the disease in many cases.

The spleen is enlarged in all my cases but it is not necessarily very large or very hard. In some of the worst cases it did not extend beyond 3" below the costal arch in most of the cases the spleen is moderately enlarged. In some very large and hard spleens I did not find any Leishman-Donovan bodies. I have often been led to suspect the disease when I found moderately enlarged spleen with extreme emaciation, cachexia and ædema of the lower extremities, symptoms which we rarely meet with in cases of malaria with moderately enlarged spleen. The liver is enlarged in a large proportion of cases. Sometimes the patients complain of pain over the splenic region which in some cases at least might be due to infarcts, often seen in this disease.

Complications—(1) Pneumonia, (2) Phthisis, (3) Dysentery, (4) Diarrhœa, (5) Cancium oris, (6) Hæmorrhage from the gums or under the skin in advanced cases, (7) Delirium or coma in some cases, shortly before death, (8) Albumen in the urine in a few cases, which might be due to a

concomitant kidney disease. I have not met with a case in which there was perforation of the bowels as has been noted by some observers

Diagnosis -The surest sign is the presence of the Leishman Donovan bodies in the blood of the spleen Spleen puncture is a simple operation and I have not met with a single accident. In some cases the patient complain of pain and tenderness for a day or two at the point of puncture Rarely there is a more general tenderness I have been more fortunate than some other observers, who have met with fatal cases of hamorrhage after spleen punc ture In one of my cases with hemorrhage from the gums hæmorrhage under the skin and marked ædema of the lower extremities spleen puncture was not accompanied with any untoward results except that the patient complained of pain for 3 or 4 days Hæmorrhage to n great extent depends I think on the size of the needle used I use the long needles supplied by B W & Co for intra muscular injections I puncture the spleen at n part in intimate contact with the abdominal wall press with a finger at the punctured spot for about 5 minutes and immediately put the patient on calcium chloride or turpentine. In bad cases I put the patient on calcium chloride for 2 days before spleen puncture One or two drops of blood drawn from the spleen are enough to demonstrate the presence of the parasites if they are present. While therefore admitting the possibility of the danger of spleen puncture as pointed out by other observers I think we may reduce it to a minimum if we do not use large needles and draw not more than a few drops of blood

I have observed the parasites post morten in the splenic veins the liver the portal veins and the bone marrow I have never found them in the superior or inferior vena cava In one case I found the Leishman Donovan bodies in the blood inside the heart within a leucocyte as will be seen in the accompanying Diagram. The L.D. bodies are very rarely found inside the red blood corpuscles. Dr. Christophers observes that he never saw unmistakable forms in red cells either in peripheral or splenic blood. In one case, however, I have observed the bodies inside red cells in the splenic blood, apparently indicating that the parasites may infect the red blood corpuscles like the malarial parasites.

All cases of enlarged spleen met with in India may not be due to Leishman-Donovan disease or malaria. Some of them may be due to some other micro-organism. Others may be due to a hitherto unknown phase of Leishmania donovani

Major Rogers lays great stress on leucocyte count as a method of diagnosis in this disease

Double remittent pyrexia and double quotidian pyrexia with double or single intermission in 24 hours, have been considered very suspicious of this disease but 1 do not consider them to be absolutely pathognomonic. On the other hand we may say that a fever which is at first intermittent in type and shows from time to time double rise with single or double intermissions in 24 hours or passes into a remittent type which also tends to show double rise, and is spontaneously attended with variable periods of apyrexia and associated with enlargement of the spleen, general wasting and cedema of the lower extremities should be regarded as very suspicious of the disease due to Leishman-Donovan bodies

In the Scientific Memoirs for 1905 Captain James lays stress upon the following group of signs and symptoms as very significant of true malarial cachexia: (1) Enlargement

^{*} It has now been conclusively proved that Leishman Donovan bodies never infect the red blood corpuscles and that the so called leishmania infected red cells are mere artifacts -Ed

of spleen (2) a temperature curve which shows definite pyrexial and apyrexial periods and (3) absence of serious symptoms throughout the period during which the condition lasts but especially so during the apyrexial intervals. Cases of Leishman Donovan disease in which the temperature curve shows pyrexial and apyrexial periods are not un common. In some of my cases, I have noticed periods of apyrexial lasting for several days. The temperature Charts were very carefully drawn the temperature being taken invariably every 4 hours and in some cases even every 2 hours. On the other hand there are many undoubted cases of true malarial cachexia in which the condition becomes progressively worse if untreated. I cannot therefore regard the above group of symptoms as absolutely pathognomonic of malarial cachexia.

Prognosis is always grave A large number of my cases died Some left the hospital in a worse condition than at the time of admission. None of my cases was benefited by treatment. Perhaps some do get spontaneously cured It is impossible in the present state of our knowledge to give a prognosis of the very early cases. Captain James believes that the tendency of true malaria is to cradicate itself from the system instead of a cachexia setting in. He holds that immunity to malaria may be acquired by repeated attacks. There can be little doubt that some malarial cases do get progressively worse without acquiring immunity. Thus it would appear that the prognosis of many cases of true malarial cachexia may be unfavourable unless properly treated.

Freatment —No drug can check the progress of this disease. I have tried quinine in small and very large doses by the mouth or by hypodermic injection without any good result. In some cases it temporanly reduces the temperature to a slight extent. Fluorides and arsenic have been tried without any effect. Cacodylate of sodium

and arrhenal seemed to do some good, though only temporarily The latter drug was also injected in ½ gr. doses into the spleen. Methylene blue, hetol and nuclein have been tried without success. In two cases I injected methylene blue into the spleen without any untoward results. An intercurrent attack of cancrum ons or a large abscess may lead to a cure.

TRANSACTIONS OF THE CALCUTTA MEDICAL CLUB

(ABSTRACT)

On the 24th June 1906 Dr Upendrannth Brahmachari read a paper on A Contribution to the Study of Fevers due to Leishman Donovan bodies (vide the Calcutta Medical Journal October 1906) Dr Kailaschandra Bose presided

Dr Satvasaran Chakravarti said that although quinine test was believed to be infallible in cases of malaria by many authorities cases sometimes cropped up which showed mala rial parasites but which did not yield to quinine. One such case was lately published in the I M G

There are cases which clinically are indistinguishable from malaria and in some cases there may be true mixed infections -although authorities differ on this point Diagnosis rests on spleen puncture and demonstrating Leishman Donovan bodies in the splenic blood and although the reader of the paper is very fortunate this little operation has sometimes been attended with death Spleen puncture is not safe specially in a country where spontaneous rupture of the spleen is not rare. Therefore, would it not be wise to trust to peripheral blood counting in at least those cases where spleen puncture would seem at all dangerous? desirable when the operation is done to take such precautions as giving calcium chloride beforehand and determining the coagulability of the blood in each case. As regards the operation itself perfect asepsis straight puncture without side movement pressure after the operation and rest in bed are better enjoined Everything I venture to say does not depend on the size of the needle Of course the finer the needle the less the chance of hæmorrhage but more the

difficulty of getting sufficient blood A larger amount is necessary when one wants to culture the organism anatomical state of the spleen to be punctured should be taken into account as well In a suspicious case, a peripheral blood count when it shows a diminution both in the red and white corpuscles, marked diminution in the number of the white blood corpuscles, especially of the polymorphonuclear variety, together with an increase in the large mononuclear and the lymphocytes, the case is decidedly associated with Leishman-Donovan bodies in the splenic blood This should be a sufficient guide in diagnosis when spleen puncture would seem dangerous The cure of cachexial fever after cancrum ons attributed to the development of staphylococcus toxin is Any marked leucocytosis brought on by any means tends to cure this disease. How often cultivators from villages suffering from other complaints are seen in the wards with huge hard spleens and many blister or actual cautery marks over the abdominal wall who are sufficiently healthy to be able to do their work They are fairly robust and free from fever These I take to be, cases of cachexial fever cured by the leucocytosis brought about by repeated cautery or blister About the prognosis of the condition though it has been said that the cases are invariably fatal, there are authorities who hold that it is not necessarily always According to the latter, if the patient eats well and has a good digestion and if he can be kept going on for 18 months he recovers

As Dr Brahmacharı says, other forms of enlarged spleen may hereafter be shown to be due to other organisms or to some unknown phase of the parasite other than the Leishman-Donovan bodies. The presence of Leishman-Donovan bodies has been demonstrated in the peripheral blood by Donovan though not found by others. As regards treatment, though it is not very encouraging, it is not so gloomy as Dr Brahmacharı would lead one to believe. Quinine though useless may prove to be valuable after the anæmia has been suitably treated for some time but certainly local puncture of the splenic region and injection of quinine have been found to be useful in a certain number of cases in which the organism was found. In laboratory animals experimental inoculation of blood containing trypanosoma has failed to infect them, if the animals were previously treated with trypan red or malachite green

Horses suffering from trypanosoma infection have been cured by these drugs according to Koch These might in suitable doses he useful in man

Owing to the lateness of the hour the meeting was adjourned

The adjourned meeting was held in the premises of the club on Saturday the 26th June Dr Kailaschandra Bose President in the chair

The further discussion on Dr Brahmachari's paper was resumed

Dr Rajendralal Dev spoke of his experience about this disease. All the cases treated by him ended fatally. He

suggested the use of X ray

Dr Haridhan Datt , thought that the early symptoms simulated those of typhoid fever The type of the disease with no fever and sudden enlargement of the spleen was very rare A slight evening rise of the temperature may not be noticed by the patient Quinine was useless but might help in diagnosis The different forms of pyrexia alluded to in the paper made the diagnosis more difficult He did not believe that a good many cases diagnosed as remittent fever were really cases of Leishman Donovan disease He thought that most of the cases seen in Calcutta were im ported from outside He could not explain why the temperature should be so variable. He objected to spleen puncture for diagnostic purposes and thought that practi tioners were not justified to have recourse to untried methods in private practice. He did not believe in the recovery of early cases or in spontaneous cure He has tried guaiacol in some cases and found it to be useful. The theory of the propagation of the disease by bugs was untenable

Dr Sashibhushan Mukherii spoke about the difficulty in diagnosis He advocated a change of climate and drugs to

improve the general health

Dr K G Sircar thought that failure of quinine in check ing fever did not imply that the fever was not malarial He referred to Dr Crombie's remarks about unclassified malarial fevers of India He related a case in which repeated blood examination showed nothing. The patient was a Eurasian adult, male residing in Calcutta. He first saw the patient after a month s slow fever with an enlarged spleen Large doses of quinine had no effect. A change to Darjeel ing did him no good and the patient came back to Calcutta

much worse A prolonged treatment with small doses of quinine, arsenic and iodine, with occasional cacodylate injections cured him

Dr Purnachandra Nandi thought that quinine might be

efficacious if combined with iron

Dr Balaichandra Sen mentioned a case in which no positive result was obtained by a blood examination. He thought that quinine would be useless in those cases where the liver was enlarged. Acid treatment might be efficacious

Dr Kedarnath Das thought that prognosis would be better if we could bring about increased leucocytosis by some means

Dr Upendranath Brahmachari, in reply, said that the failure of quinine in malaria (proved by a blood examination) is almost unknown. Such cases were generally complicated with the other disease, thus explaining the failure of quinine test. He did not think that leucocyte count, as helping diagnosis, was always satisfactory. As complications were common in Leishman-Donovan disease, the result of leucocyte count was bound to be vitiated. In children leucocyte count was always doubtful

The president observed that the effect of guinine depended on the way it was exhibited. Quinine test was not always diagnostic. The mere presence of malarial parasites in the blood did not indicate that Leishman-Donovan bodies were absent. Both may exist together. He did not agree with the lecturer that spleen puncture was safe.

SPORADIC KALA AZAR IN CALCUTTA WITH NOTES OF A CASE TREATED WITH ATOXYL

Kala azar as used in this article may be defined as the disease caused by the Leishman Donovan bodies lts epidemic manifestation which is seen in Assam more frequently goes by this name. It is endemic round about Calcutta and probably occurs in the city itself

Last year I examined the spleen blood of nearly 150 cases of enlarged spleen admitted into my wards in the Campbell Ho pital and found the parasite of kala azar in 60 cases. From this it is seen that cases of this disease are frequently admitted into the Calcutth hospitals.

No two cases came from the same house and I have not succeeded in corroborating the fact that the disease is limited to individual houses or families 91 7 per cent of my cases were Hindus while 8 3 were Mahomedans giving a proportion of nearly 11 to 1. Out of the total number of patients admitted into my wards in 1907 about 75 per cent were Hindus and 17 5 per cent Mahomedans giving a proportion of nearly 4 to 1. It will thus be seen that Hindus are more frequently affected by the disease than the Mahomedans and if this fact is corroborated by more extended observations it may probably throw some light on the etiology of the disease. Thirty three per cent of my cases were below the age of 20. I have not met with any case above the age of 35. The cases came to my wards during all the parts of the year, the largest number of admissions being

during the period of April to August, but especially in the month of May

It was difficult to trace from the history of the cases where the disease was contracted. Many of them pointed, however, to the disease having been contracted in the neighbourhood of Calcuita, and a few seemed to have contracted the disease in the city itself, as they never went out of it None of my cases came from the epidemic area of Assam Some came from Orissa, Eastern Bengal, district of Murshidabad, and Behar

Most of my cases were chronic, with history of illness for several months, and with the spleen extending 3 inches or more below the costal arch. In a large majority the liver was also moderately enlarged, while in a very few cases the liver was very much more enlarged than the spleen, which extended just an inch or so below the ribs. I have not met with a case in which the spleen was not at all enlarged

SYMPTOMS

As regards the early symptoms of the disease our knowledge is still very limited. The patients met with in the hospital wards have suffered for a long time. I give here the summary of the early symptoms of the disease as gathered from the patients. (Vide Indian Medical Gazette, Vol. XLI, January, 1906.)

In one class of cases we get a history of intermittent attacks of fever, quotidian for some time, and later on becoming irregular with or without rigors, lasting for several months and not benefited or partly benefited by quinine ln a second class there is a history of a few previous attacks of fever, continuing for some time on each occasion, the spleen becoming slowly and steadily enlarged. In a third class there is history of attacks of low fever, continued for several months, with a progressive enlargement of the spleen not benefited by any treatment. In a fourth class

there is a history of one or two attacks of remittent fever more or less resembling typhoid in a fifth class the patient gives a history of gastro intestinal troubles with dysenteric or diarrhoea attacks followed by cedema of the lower extremettes and attended from time to time with ague like attacks of fever in all these cases the patient invariably says that there was very slight or no benefit from quinine Possibly a large majority of cases begin with attacks of remittent fever

Among the other symptoms that I have observed may be mentioned progressive emaciation anæmia cachexia cedema of the extremeties diarrheea dysentery which may sometimes be very obstinate and hæmorrhage from various parts of the body such as the skin and the mucous membranes

Œdema of the extremities is sometimes an early symp tom On the other hand it may be absent even up to the last It is not easy to explain the cause of this cedema It appears that we can distinguish two classes of cases one in which cedema is a marked symptom and the other in which it is not

I have elsewhere described in detail the various types of pyrexia that I have observed in this disease (Indian Medical Gazette January 1906) Besides these, there may some times be pyrexia of the pyeemic type there being more than two remissions or intermissions in twenty four hours. Rarely the disease takes on an apyrclic course for an indefinite length of time.

BLOOD COUNT

In the main my observations corroborate those of Major Rogers (Lancet March 2nd 1907) Leukopenia is extreme and in many cases the proportion of white to red cells is less than 1 1 000 (Table 1)

TABLE I

	R ed Corpuscles	White Corpuscles	Proportion
1	2,390,000	1,720	1/1389
2	2,530,000	2,530	1/1000
3	2,740 000	1,792	1/1529
4	3 440,000	2,386	1 / 1441
5	3,170,000	2,380	1/1332
6	2,660,000	1,160	1/2293
7	2,760,000	1,527	1/1807
8	2,920 000	1,320	1 2212
9	2,120,000	1,736	1/1221
10	2,330,000	1,500	1/1456

In a number of cases, though there was a marked leukopenia, yet the proportion of white to red was higher than $\frac{1}{1000}$ There are probably very advanced cases with extreme anæmia (Table II)

TABLE II

	Red Corpuscles	White Corpuscles	Proportion
1	1,400,000	2,170	1/645
2	860 000	2,250	1, 382
3	1,480 000	3,870	1,382
4	1 300 000	3,300	1/383

In many cases complications often lead to increase in the leucocyte count (Table III)

TABLE III

	RdC p 1	Wht Cpls	Pptn	C mpl t
1	2 880 000	7 662	1/376	Pn um
2	896 000	2 768	1/323	C um
3	3 070 000	3 500	t 817	24 hu ft i pl t f l g bl t o th pl g

Lastly cases that have been recovering or have recovered from cancrum ons or have been treated with atoxyl for some time may give a higher relative leucocyte count than 1/1 000 (Table IV)

TABLE IV

	RdC pul	Wht Cp 1	Ppiln	C mpl t
1 2 3	2 340 000 2 600 000 3 440 000	4 569 3 340 5 060	1,512 1/778 1/529	R y f m T tm nt wth

Comparing the leucocyte count in kala azar with what we observe in malarial cachexia exactly the same conclusion as that of Major Rogers is reached namely that a ratio of red to white corpuscles below 1/1 500 is almost diagnostic of kala azar but as we have shown above we may meet with cases without any apparent complication or some with well marked complications or others that have taken a favourable turn in which the proportion may be higher than 1/1 000

I append here a table showing the blood count in malarial cachexia in which the proportion of white to red is much greater than 1/1 000 (Table V)

TABLE V

Malarial Cachexia

!	Red Corpuscles	White Corpuscles	Proportion
1	3,240,000	8,000	1/405
2	2,208,000	3,372	1,655
3	2,384,000	4,863	1/490

COMPLICATIONS

(1) Pneumonia, (2) phthisis, (3) dysentery, (4) diarrhoea, (5) cancium ois and its attendant complications, (6) hæmorrhage from the gums or under the skin in advanced cases, (7) delirium or coma, in some cases, shortly before death; (8) albumen in the urine in a few cases, which might be due to concomitant kidney disease, (9) paraphimosis, (10) hæmoptysis, (11) hæmatemesis, (12) melæna, (13) epistaxis, (14) cedema of the extremeties, (15) splenalgia due to infarcts in the spleen, (16) hæmorrhoids, which may bleed obstinately and profusely. (17) large abcesses. I have not met with a case in which there was perforation of the bowels, which Captain Christophers describes in some of his cases.

Prognosis

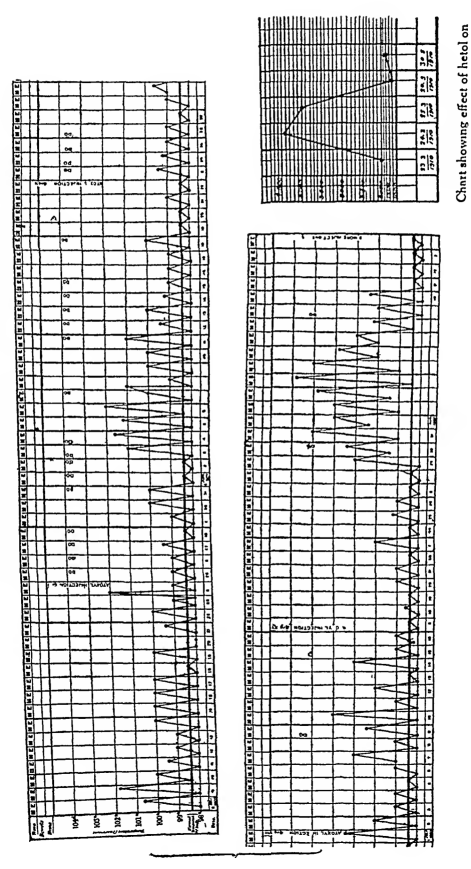
According to my observations, the prognosis is always grave—I have not the record of a single—case which I could pronounce cured, 40 per cent—of my cases died in the hospital, the most—frequent—cause of death being intractable diarrhœa or dysentery

TREATMENT

So far as my observations go, no drug can kill the parasites I have used the following drugs without success

leucocyte count

Temperature chart of a kala azar case treated with atoxyl (Para 3)



(1) quinine internally (2) quinine hypodermically (3) fluorides (4) arsenic and some of its new preparations such as sodi cacodylate arrhenal and atoxyl in some cases arrhenal and sodi cacodylate were given hypodermically (5) methylene blue internally (6) methylene blue hypodermically (7) izal in increasing doses (8) cyllin in increasing doses up to half a drachm thrice a day

Encouraged by the results of the use of atoxyl in trypano somiasis I gave it an extensive trial in a few cases one of which has been under my observation for nearly eight months and under atoxyl treatment for nearly six months. As far as I am aware there are no records of a single case in which atoxyl was used for such a prolonged period in kala azar.

The patient set 30 a Hindu had slight sedema of the lower extremities at the time of admission, the spleen ex tending 8 in from the left nipple to the middle line

The effects of the treatment were as follows

1 Body Weight-

		٠.	••
May 25	1907	6	0
August 12	1907	7	4
September 9	1907	7	51
November 16	1907	8	0
December 29	1907	8	51

C. 11.

2 Blood Count—

Dι	RdC pu t	Wht C pu t	Ppt
M y 2 1907	2 769 000	1 527	1/1807
A g t 16 1907	2 500 000	1 600	1/1562
Spt mb 7 1907	2 280 000	1 950	1/1170
N mb 18 1907	2 600 000	3 340	1/778
J y 7 1908	3 449 000	6 500	1/529

- 3 Pyrexia—(There is some effect on the Temperature curve)
- 4 General condition—The patient looks much better in health, the ædema of the extremities has completely disappeared. He is stouter than before, is less anæmic, is not cachectic and has got better appetite.
- 5 Toxic action—The drug was almost non-irritating Except on one occasion there was very slight local irritation. On this occasion there was a slough which I think was due to the solution being too hot—In other cases in which I have given injection of atoxyl no untoward—local—symptoms were met with
- 6 Effect on the parasites—They are still found in the splenic blood Some of them present a granular appearance, but they are still to be seen in large numbers
 - 7 Spleen—Not much diminished in size

Before concluding, I would just mention one point, that spleen puncture, though it has been pronounced dangerous by the highest authorities, has never led to any single accident in any of my cases. I have found that a large hypodermic needle is sufficient to enable one to draw one or two drops of blood, which are almost always sufficient to show the parasites if they are present. I would therefore recommend that not more than one or two drops of blood should be drawn from the spleen for ordinary diagnostic purposes. It appears to me that the danger of spleen puncture may be reduced to a minimum if one uses a hypodermic needle, gives the patient calcium chloride before and after puncture, enjoins absolute rest in bed, and puts pressure upon the punctured spot for about fifteen minutes after the operation.

Conclusion

1. Sporadic kala-azar is frequently seen in Calcutta hospitals and is probably endemic in Calcutta itself. Hindus are probably more affected than Mahomedans.

- 2 The leucocyte count is of great diagnostic help but not absolutely so in kala azar
- 3 Atoxyl is borne in very large doses (grs xv in one injection) by kala azar patients—It is perfectly harmless and non irritating—Its effects on the parasites is perhaps very slight—While improving the patient s—health—it does not remove the cause—though it has apparently some effect upon the pyrexia—In order that it may do any good it must be given—in very large doses (grs—xv) by injection—every seven or ten days—continued for several months
- 4 Spleen puncture—It is practically safe if one uses a hypodermic needle and does not draw more than one or two drops of blood *

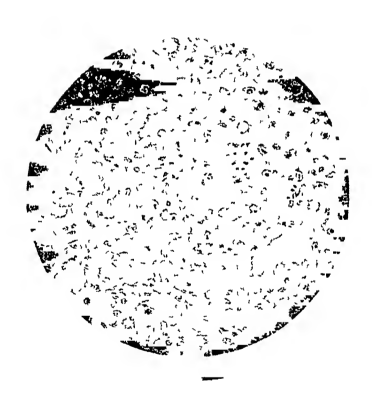
FATTY LIVER IN KALA-AZAR

In the British Medical Journal, May 30, 1903, I pointed out that in cases of Fala-azar, the liver is frequently enlarged. This enlargement is in some cases due to an intracellular circhosis, as pointed out by Major Rogers. In one case I observed extensive fatty changes in the liver, which, I consider, is a very rare condition in Fala-azar, It has not, as far as I am aware, been describe I by any previous observer.

Condition on admission—The patient, at 18, was admitted into my wards with history of fever coming on irregularly for about eight months. He was markedly emaciated and cachectic. The spleen was slightly enlarged, extending about an inch below the costal arch. The liver extended about an inch and a half below the costal margin. The spleen blood showed a large number of Leisliman-Donovan bodies. The lungs and the heart showed nothing abnormal. There was no complication at the time of admission into the hospital

After history—After a fortnight he began to suffer from very obstinate diarrhoa, which terminated fatally. There was no ascites, and there were no enlarged veins in front of the abdomen nor other signs of portal obstruction. There was no cedema in the lower extremities. The fever was irregular, sometimes presenting the double remittent type and sometimes hectic. The patient died in the hospital twenty-five days after admission. He was treated with injections of atoxyl. A few days before death he suffered from an abscess in the right arm, and also an inflammatory swelling in the left.

[Reprinted from the British Medical Journal, Vol 11, September 26, 1908]



Section of liver showing extensive fatty degeneration of the cells, more about the peripheral portions of the lobules of the liver (Vide para 3)

Blood Count—Red corpuscles 1 957 000 white 2 723 proportion, 1 in 720

Post mortem Examination—The body was emaciated subcutaneous fat diminished heart valves healthy lungs nothing abnormal

The liver was enlarged smooth and yellows him colour lts margin was somewhat rounded on section it had a yellow mottled appearance When the surface was crap droplets of fat could be seen on the knife Micro copicall, (see Diagram) extensive fatty degeneration of the liver cells was noticed rather more about the peripheral portions of the lobules of the liver In many places the liver cells were shrivelled and many contained small oil globules scattered throughout their protoplasm In a few of the liver cells there could be seen deposits of granules of golden brown pigment resembling hæmosiderin. In some of the lobules there was a very marked engorgement of blood in the centre (not shown in the Diagram) In some places there was a slight formation of new bile ducts There was all o a growth of delicate connective tissue appearing between the liver cells

Reference

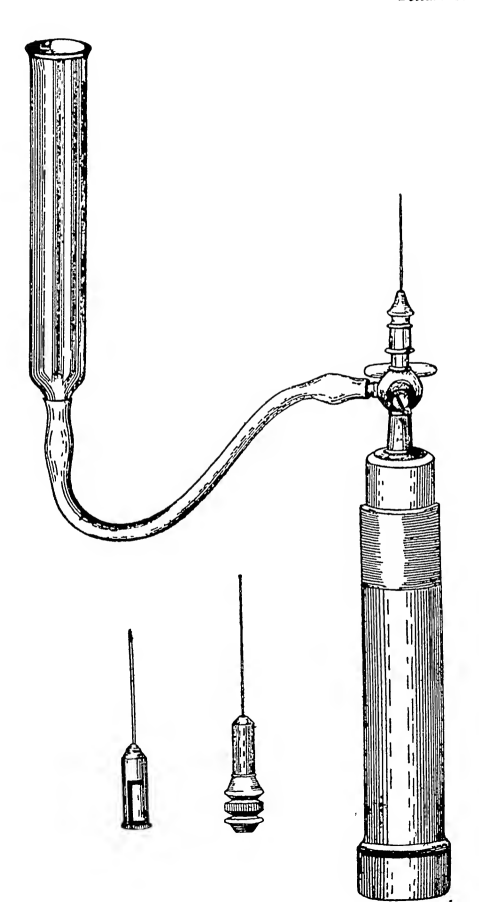
I L ROGERS (1908) F th T p

A PRELIMINARY REPORT ON THE TREAT-MENT OF KALA-AZAR WITH INTRA-VENOUS INJECTION OF METALLIC ANTIMONY

The use of metallic antimony in a state of finest subdivision in the treatment of kala-azar has not been noted by any previous observer. In a disease like kala-azar in which the parasites reside in such organs as the spleen and the bone marrow and in which very few parasites are found in the peripheral circulation, the use of the soluble salts of antimony may be followed by such quick elimination of the drug that they may be excreted before exerting any marked influence on the parasites in the spleen and the bone marrow, while metallic antimony may be retained for a much longer time if introduced into the circulation the treatment of the allied disease of trypanosomiasis by intravenous injection of tartar emetic or other soluble salts of antimony, it has been found that though they quickly free the peripheral blood from typanosomes, still the parasites remain in the internal organs So far, therefore, the treatment of trypanosomiasis by means of intravenous injection of soluble salts of antimony has not been a permanent success In kala-azar, in which the parasites live mostly in the internal organs, the same line of treatment may, from these theoretical considerations, be not expected to do much good However, the treatment of kala-azar with tartar emetic and other soluble salts of antimony will form the subject of a future communication, though it may be remarked here, in passing, that good results with tartar emetic

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have been recently reported by several observers. The present paper is a preliminary report but the results obtained so far certainly justify further investigations.

The method of administration is the author's own and is described as follows —

An all glass 10 c c syringe is fitted with a three way stop cock the remaining two ends of which are fitted to a platinum needle and a rubber tubing attached to the nozzle of a piece of glass tube respectively. A stout hypodermic needle with a specially constructed blunt canula inside may be used in place of an ordinary one. This canula when pushed through the needle will prevent the puncture of the vein a second time during the process of injection of the metallic antimony (see Figure). One grain of metallic antimony is made into a thoroughly homogeneous paste with sufficient liquid glucose in a glass morter and then mixed with 20 c c normal saline. (The glucose added is just sufficient to make a 5 per cent solution with 20 c c of normal saline.)

The stop cock is so arranged that the syringe may be made to communicate alternately with the needle or the glass tube by turning the stop cock Half the suspension is now sucked into the glass syringe and after being freed from any bubbles of air it is injected into a vein The glass tube is now filled with a of the forearm portion of the remaining suspension which is sucked into the glass syringe after freeing the rubber tubing of any air bubbles and then the suspension is again injected In this way the whole of the suspension is injected into the vein Any sediment of antimony left inside the syringe is subse quently mixed with normal saline containing 5 per cent glucose and then injected into the vein. This process is repeated several times till no antimony is left inside the syringe About 40 to 45 c c of normal saline is required to mject I to 1 grs of metallic antimony In all the cases

recorded below, the diagnosis was made by the presence of Leishman-Donovan bodies in the splenic blood

The first two cases were at first treated with tartar emetic and Plimmer's salt (sodium antimonyl tartrate) Then after some days they were treated with intravenous injection of metallic antimony

Case No 4 was treated with intravenous injection of metallic antimony and electrical. This patient had marked jaundice with cedema of the extremities, and there was a large amount of albumen in the urine. These completely disappeared during treatment.

Case No 5 had cancrum orts on admission which disappeared after treatment

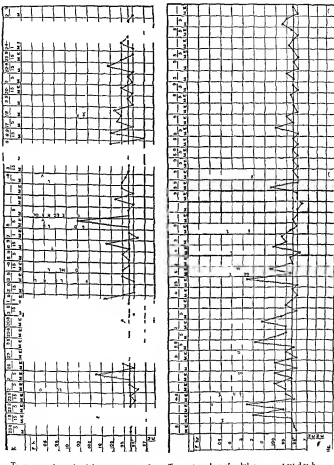
The results so far obtained in these cases are recorded in the accompanying tables and the effects on the temperature are shown in the accompanying Charts.

It will be seen from the above that so far the results are encouraging and justify further investigation. No untoward results, such as plugging of the capillaries, have followed the intravenous injection of metallic antimony. Sometimes the patient suffered from rather severe diarrhæa, which, however, stopped in 24 to 48 hours.

The advantages of intravenous injection of metallic antimony given in the above way are as follows —

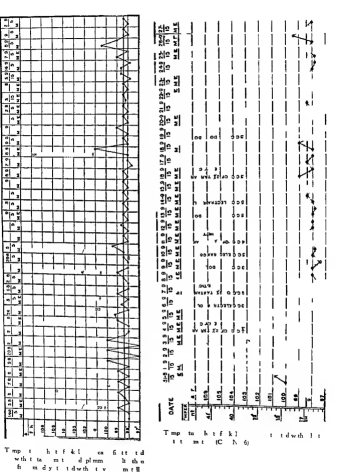
- (1) A small quantity of normal saline is required for injecting the antimony. This is an advantage, as kala-azar patients are frequently liable to suffer from high fever and rigors after injection of large quantities of normal saline.
- (2) An ordinary hypodermic needle may be used for the purpose, if it fits the stop-cock
- (3) The glucose is added to raise the specific gravity of the suspending fluid, and thus it reduces the settling down of the metallic antimony inside the injecting syringe to a minimum

Case No 6 was treated with electargol and tartar emetic and is recorded here for comparison



Temp t hat f klaza c t td wth lect g l d m tll ntm ny C No 4) Temp to h tof kal-a teatdwh trt mat antm nyl d mat t t nd m tlle tem ny tP 2)





antm y (P

2)



TABLES SHOWING THE EFFECTS OF TREATMENT

Case 1-Sitaram

	{	Size of	Spleen	
Di	Body w ght	Lwrb d xtnd lngth litnppl	Lng t tnse dm t bg fm	Bl d x m n t
3 8 15	3 t 5 li	5 blw ctl	let { f m d { dl	RBC 2 000 000 WBC -2 400
19815	3 1 31 16	D	Do {	R B C -2 400 000 W B C -2 400 b - 38
27 8 15	D	Do	D {	RBC = 2 400 000 WBC = 3 000 Hb = 38
6 9 15	D	41 D	2 Do {	RBC 2600 000 WBC 6400 Hb 44
15 9 15	Do	Do	Do	RBC -3 600 000 WBC -7 200 Hb -569
27 9 15	Do .	3 D	21 D	RBC -4 000 000 WBC 8 400 Hb - £0

Case II-Ramptan

		SIZE OF	SPLEEN	
D t	Body Lw bo d t d l g th I ft n ppl		Long t t n se d m te b g ns f m	Blood m t
14-8 15	3 t 4lb	41 blw	I ght { f m d } d!]	R B C 2 800 000 W B C 5 600 Hb 44%
20 8 15	3 t 6 lb	Do	D	RBC -2 600 000 NBC -5 600 Hb -46%
3 9 15	Do	Do	a lft of md dle le	RBC-2400 000 WBC-4800 HI-48
18-9 15	3 : 815	Do	3 lft f md	RBC -3 200 000 WBC -8 800 Hb -58%

Case III-Mati

		Size of	SPIFFN	
Date	Body weight	Lower bor der extends along the left nipple line	dinineter begins	Blood examination
13 9 15		5' below costal arch	2 right of mid dle line	RBC -2 000 000 WBC 2400 Hb 46,
27 9 15		3' Do	l left (of mid { dlc line {	R B C -2 800 000 W B C -3 000
1 10 15		2 Do	Do {	R B C -2 800,000 W B C -2 803 Hb -52′,

Case IV—Durgadhan

		Stze of	· Spleen	
Date	Body weight	Lower bor der extends along the left nipple line	Longest transverse diameter begins from	examination
21 8-15	5 st 2 lbs	5½″ below costal arch	2½" right { of mid dle line	RBC -3,400 000 WBC -4,000 Hb -48%
3-9 15	5 st 3 lbs	Do	D _o	R B C—3 800,000 W B C —3,200 Hb —48%
11 9 13	5 st 6 lbs	D _o	Do {	R B C -4,000 000 W B C -3,200 Hb -62;
27 9 15	Do	4" Do	≟″ Do	R B C4,200,000 W B C6,000 Hb6'%

Case V-Kası

Dt	B dy w ght	Size of Spleen	Blood x m n t on
18-9 15 27 9 15		Spleen felt jut bel w { the cotl rch Spleen cann t b f lt { below the cotlach	R B C - 2 400 mm W B C - 4 800 Hb 50 R B C - 2 800 000 W B C - 5 600
			Нь —52%

Case VI-Narayan

		Size of	SPLEEN	
Dt	Body w ight	Low rbo drtd algth lttnippl lne	Log t tr d m t beg n f m	Blood xmntn
4 9 15	4 t 6 li	5 belw	l ft of {	R B C -2 800 000 W B C -1 600 Hb -46
17 9 15		4j Do	13 0 {	RBC 2 600 000 WBC 4 400 Hb 50°
18 9 15	4 t 4j lb	4 D	2 Do {	RBC -26 0 000 WBC -4 000 Hb -54 6
2 10 15	Do	D	D ₀ {	RBC —3 800 000 WBC —6 400 Hb —707

Case III-Mati

		Size of	SPLEEN	
Date	Body weight	Lower bor der extends along the left nipple line	Longest transverse diameter begins from	Blood examination
13 9 15		5' below costal arch	2 right of mid dle line	RBC 7,000 000 WBC 2 400 Hb 46%
27 9 15		3" Do	l' left of mid dle line	R B C —2,800,000 W B C —3,000
1 10 15		2′ Do	Do {	R B C —2 800,000 W B C —2 80J Hb —52%

Case IV—Durgadhan

		Size of	SPLEEN	
Date	Body weight	Lower bor der extends along the left nipple line	Longest transverse diameter begins from	► Blood examination
21 8 15	5 st 2 lbs	5½″ below costal arch	2½″ right { of mid- dle line	RBC -3,400 000 WBC -4 000 Hb -48%
3-9 15	5 st 3 lbs	Do	D _o	R B C—3 800,000 W B C —3,200 Нь —48%
11 9 13	5 st 6 lbs	Do	Do {	R B C -4,000,000 W B C3,200 Hb62 \(\)
27 9 15	Do	4″ Do	2" Do {	RBC4,200,000 WBC6,000 Hb6'%

Case V-Kası

D:	Body wight	Size of Spleen	Blood amin ti n
18-9 15		Splen flipust blow (th cot lach	R B C -2 400 000 W B C -4 800 Hb 50
27 9 15		Spleen c nnot be felt below the c t l rch	RBC -2 800 000 WBC -5 600 Hb -52%

Case VI-Narayan

		Size of	Spleen	
D t	Body w Ight	Low bodrtder to describe the second s	L ge t tr e d m t r begin lrom	Blood mì tỉ n
4 9 15	4 t 6 11	5 below ctl rh	left of anddl	R B C -2 800 000 W B C -1 600 Hb46
11 9 15		4] Do	11 Do {	RBC -2 600 000 WBC -4 400 Hb -50
18 9 15	4 t 4 <u>1</u> lb	4 Do	2 Do {	RBC-26 0 000 WBC-4 000 Hb-54/
2 10 15	D	D _o	D {	RBC —3 800 000 WBC —6 400 Hb —70%

FURTHER OBSERVATIONS ON THE TREATMENT OF KALA-AZAR AND CASES TREATED WITH METALLIC ANTIMONY, SODIUM ANTIMONYL TARTRATE, FORMALDEHYDE AND OTHER DRUGS

SECOND REPORT

This paper is a continuation of my previous paper which was published in the *Indian Medical Gazette*, December, 1915. It includes fresh cases treated with metallic antimony and other drugs and further observations about cases already reported.

- I Cases treated with metallic antimony —
- (a) Cases already reported -
- 1 S—The patient left hospital after being free from fever for nearly a month He had altogether five injections of metallic antimony, half a grain each time

Result of blood examination —

- R B C--2,000,000 W B C --2,400 on 3rd August, 1915
- R B C —4,000,000 W B C —8,400 Hb —60% on 27th September 1915
- Size of spleen —(1) 5" below the costal margin on 3rd August, 1915 (2) 3" below the costal margin on 15th September, 1915



DAT	E	8-8 21 23	10-9 15 M.E	15	15	15	15	15	15	15	15	15	15	15	15	15	15	25 9 15 M E	15	15
TEMPER																				
Cont	Fahr				רום ו															
41	106°				OF METAL															
	105°				OR OF ANT			U.			00									
40°	104°				-101	-		OF METALL												
	103°	-0	1	₽ <u>₽</u>	Δ		R	OF ANT												
39 -	102°	1				Å		1 68			OQ									
	101°	7	1			R		1_			_{\lambda}									
38	100°		M					4	1	ĹŢ										
37°	99°		L	<u> </u>			<u> </u>				V					-8				
NORF AL	98°					-	₩_			4	-	يمطر		- 24			_A	حر	~	-
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Pulse	· {E												-3-							
Resp	{M €																			
Bowe	ls																			

Temperature chart of a kala azar case treated with three injections of metallic antimony half a grain each time (Para 1, No 2)

2 R—The patient left hospital after being free from fever for nearly a month. He had altogether three injections of metallic antimony, half a grain each time.

Result of blood examination -

- RBC-2800000 WBC-4600 Hb-44° on 4th August 1915
- R B C -- 3 800 000 W B C -- 8 800 Hb -- 66 on 29th September, 1915
- Size of spleen —(1) 43' below the costal margin on 14th August 1915 (2) 3' below the costal margin on 3rd October 1915 (3) Longest transverse diameter diminished by nearly 4
- 3 M—The patient left hospital after being free from fever for nearly three weeks. He had altogether four injections of metallic antimony.

First dose half a grain and subsequent doses one grain

Result of blood examination -

- R B C -- 2 000 000 W B C -- 2 400 Hb -- 46% on 13th September 1915
- R B C -- 3 600 000 W B C -- 4 400 Hb -- 58% on 9th October 1915
- Size of spleen —(1) 5 below the costal margin on 13th

 September 1915 (2) 1½ below the costal margin on 11th October 1915
- 4 D—The patient left hospital after being free from fever for nearly a month and a half. He had altogether five injections of metallic antimony first dose half a grain second third and fourth doses one grain each fifth dose one and a half grains.

Result of blood examination —

R B C —3,400,000, W B C —4,000 Hb —48% on 21st August, 1915

R B C -4,600,000 W B C -10,400 Hb -68% on 29th October, 1915

Size of spleen —(1) 5½" below the costal margin on 21st August, 1915 (2) Cannot be felt below the costal margin on 3rd November, 1915

5 K—The patient is still in hospital. He had a very severe attack of acute enteritis and also suffered from pleurisy for a few days. For some time his condition was very serious. As mentioned in my previous paper he had cancium oris at the time of admission. He has been free from fever for nearly a month and a half, except for a short period when he had pleurisy. He had altogether two injections of metallic antimony, first dose, half a grain and second dose, one grain

Result of blood examination —

R B C -- 2,400,000 W B C -- 4,800 Hb -- 50% on 18th September, 1915

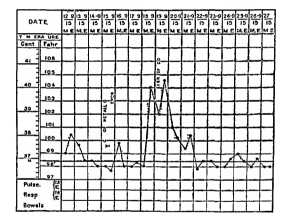
R B C -2,600,000 W B C -5,000 Hb -55% on 28th November, 1915

Size of spleen —(1) Just felt below the costal margin on 18th September, 1915 (2) Cannot be felt below the costal margin on 28th November, 1915

The general condition is improved and the patient is having a slow convalescence. The cancium oils is cured but the patient in still very feeble

(b) New cases —

Three more cases were treated with metallic antimony which are recorded here





6 L—Patient was suffering from high fever for some time before admission. He had altogether five injections first dose being half a grain second third and fourth doses one grain each and fifth dose one and a half grains (See Table 1 and temperature Chart I) The patient is doing very well

Size of spleen —(1) 4 below the costal margin on 12th
October 1915 (2) Cannot be felt below the

- 7 Putu—The patient had been in hospital for a short time and has left since
- 8 Abdul—The patient had been in hospital for a short time and his case will be reported later on

II Treatment with intravenous injection of galyl -

The brilliant results obtained with galyl in the treatment of sleeping sickness led me to try this drug in four cases Each had one injection of 25 gramme but no good results have followed its use. The drug should however be given a more extended trial, which is impossible in a hospital on account of its cost.

Ill Treatment with intravenous injection of form aldehyde —

The patient named Gopi a hospital servant was ad mitted into my wards on 11th June 1915 suffering from fever of remittent type. He had another attack a short time ago. On admission his spleen was found to extend four inches below the costal margin, and on spleen puncture a large number of Leishman Donovan bodies were found. The patient was treated with intravenous injection of formalde hyde. Three injections of 40 c.c. of a solution of formal dehyde in normal saline (1 in 4 000) were given (See temp. Chart 11.)

The patient left hospital when his spleen extended two inches below the costal margin having much improved in health. He went to his native village and has come back in

perfectly healthy condition This is one of the few cases in hospital which has been under my observation for nearly six months. His general condition is now very much improved and he is doing his work in hospital as a perfectly healthy man (See Table II for his blood condition)

IV Treatment with Plimmer's salt (sodium antimonyl taitiate)—I have already reported two cases treated with this salt which were subsequently treated with metallic antimony Four more cases have been treated with the same drug

The following are the notes on one of them -

The patient, M, came under my treatment on 1st October, with enlarged spleen extending about 4 inches below the costal arch. He was at first treated with galyl which did him no good as the temperature chart and the result of the blood examination will show. After 10 days he was put on a course of Plimmer's salt. The results of blood examination and the temperature Chart are appended herewith (Chart III and Table III.)

The results seem to be, so far, similar to what has been reported in cases treated with tarrar emetic. The doses are similar to what I generally follow in the case of tarrar emetic. The patient seems to have much improved in health. The salt used by me was perfectly pure, having been specially prepared for my work. I have given Rai H. N. Ghose Bahadur some of this pure salt to use in his wards.

V Treatment with intravenous injection of tailar emetic and sodium antimonyl tartrate — Three cases were treated with both the drugs, the idea being that when the higher doses were used, the sodium salt was preferable. One of these patients, a girl named K, æt 11, has remarkably improved under this treatment. (See Chart IV and Table IV.)

Another case, a boy, æt 11, was given from ½ c c to 4 c c of 2 per cent solution of tartar emetic for 7 days and then he was treated with 5 c c to 6 c c of 2 per cent solution

¢

of the sodium salt every two days for some time and afterwards every four days. He was altogether given 14 in jections. The result has been that the spleen has gone down from 4 to ½ below the costal margin and the temperature has come down to normal. The R.B.C. W.B.C. and Hb. have gone up from 4400 000 2 910 and 35° before treatment to 5 600 000 9 100 and 55° respectively after treatment in n. month s time.

The highest dose of the sodium salt used for nn adult was 7 c c of 2 per cent solution

1'1 Treatment with tortor emetic and other drugs combined—In one case the patient was put on Lerberine sulph (21 grs) and nuclein capsules thrice a day for two months and a half along with tartar emetic given intravenously. The result so far obtained does not seem to be different from what has been obtained from tartar emetic alone. In another case berberine sulph has been und intravenously for some time. In one case gally was administered ofter a course of treatment with tartar emetic but this did not influence the course of the disease.

Retention of antimony in the tissues when introduced into the system as nietalke ontimony—la my last paper. I pointed out the possibility of the retention of natimony in the tissues such as the spleen for a much longer time when introduced into the circulation as metallic antimony than when introduced in the form of soluble salts. The following investigations have confirmed this view—A patient treated with tartar emetic died of dysentery 24 hours after the last injection no trace of antimony could be detected in the spleen and the liver by chemical examination. Another patient who had an injection of half a grain of metallic antimony also died of dysentery 132 hours after the injection and traces of antimony could still be detected in the spleen and the liver by chemical examination thus showing that antimony introduced into the circulation as

metallic antimony is retained in the spleen and the liver for a much longer time than when introduced as a soluble salt

I am indebted to Rai Chumlal Bose Bahadur, Chemical Examiner to the Government of Bengal, for the above chemical analyses

Conclusions

- (1) So far as the above cases go to prove, metallic antimony seems to produce very marked beneficial effects in kala-azar, and the effects tend to be permanent
- (2) The soluble salts of antimony, such as tartar emetic and sodium antimonyl tartrate, are also very beneficial in the treatment of the disease.
- (3) Metallic antimony introduced into the circulation remains in the spleen and the liver for a much longer period than when introduced in the form of soluble salts. It may, therefore, be expected to give rise to more marked and permanent results. Not more than five injections have been given to any of my cases. It appears that not more than three or four injections are required to bring about what appears to me a permanent cure
- (4) The results obtained from combining antimony treatment with other drugs such as galyl, berberine sulph and nuclein, do not, so far, seem to differ from what follows the treatment with tartar emetic itself
- (5) Galyl has been tried in four cases without any effect, but evidently it must be given a further trial before its effect can be determined
- (6) One remarkable case of recovery has been recorded following intravenous injection of formaldehyde
- (7) Attempts are being made to prepare a colloidal solution of the metallic antimony, and if this succeeds, the colloidal solution of the metal will perhaps be the ideal preparation of antimony to be adopted in the treatment of kala-azar

A portion of this paper was read before the Calcutta Medical Club last November and some of the cases were mentioned in the meeting of the Medical Section of the Asiatic Society of Bengal last December

TABLE I

L —Patient treated with metallic antimony

D te	S z of spleen	Rult of blood mnat
13 10-15	4 blowth cot 1 ch	RBC-2 140 000 WBC-2 600 Hb-32%
23 10 15	l do	{RBC -2 500 000 {WBC -2 100
1 11 15	Cann t b felt bel w tl	RBC-2200000 WBC-4000 Hb-32%

TABLE 11
Gopi—Patient treated with formaldehyde

D t	Sz of ple n	R It fblod m t
12 6 15	4 b low th t I m gin	RBC-3800000 WBC-3200
5 7 15	2 d	RBC-4200000 WBC-4400 Hb-54/
1 11 15	Cn ot be flt belw the talm gm	RBC -5 000 000 WBC -11 200 Hb -859

TABLE III

M.—Patient treated with Plimmer's salt (sodium antimonyl tartrate)

Date	Sız	e of spleen	Result of blood examination
7-10-15	4" below the costal arch		(RBC-2,400,009 {WBC-1,200 Hb-38%
13 10 15	4"	do	{ R B C1,809,000 W B C2,000 Нь30%
27-10-15	2"	do	RBC -2,600,000 WBC -3,600 Hb40%
9-11-15	1"	do	RBC -3,200,000 WBC -6,000 Hb -55%

TABLE IV

Patient treated with tartar emetic and sodium antimonyl tartrate

Date	Body weight	Size of spleen	Result of blood examination
26 10-15	2 st 10 lbs	4" below the costal arch	{ R B C -2,950,000 { W B C -3,200
20-11-15	1	3" do	RBC -2,800,000 WBC -2,400 Hb -38%
21-11-15		2" do	RBC -2,900,000 WBC -5,200 Hb - 42%
3 12-15	3 st 2 lbs	Cannot be felt below the costal arch	RBC 3,000,000 WBC 5,600 Hb 44%
15 12-15		do	RBC3,900,000 WBC10,600 Hb60%

THIRD REPORT ON THE TREATMENT OF KALA AZAR WITH SPECIAL REFERENCE TO THE USE OF ANTIMONY AND FORMALDEHYDE*

This paper is a continuation of my papers on the treatment of kala azar published in the Indian Medical Ga.ette last December and January. It includes fresh cases and further observations about cases already reported

The paper can be conveniently divided into the following parts -

- 1 Cases treated with metallic antimony
- 2 Cases treated with compounds of antimony
- 3 Cases treated with intravenous injections of antiseptics, e.g., formaldehyde, eusol
 - 4 Alkaloidal therapy
 - 1 Cases treated with Metallic Antinony

(a) Colloidal Metallie Antimony

In my last communication in the Indian Medical Galette
I pointed out that colloidal metallic antimony would perhaps
be the ideal preparation of antimony to be used in the treat
ment of kala azar

The use of colloidal metallic antimony in the treatment of kala azar has not been noted by any previous observer Reference to its use in therapeutics is very meagre. The following cases are therefore of much interest as being the

R ad before the M d cal Section of the A t tie Society of Bengal on 26t! April 1916

first recorded cases showing marked benefit by the use of colloidal antimony

The patient Bhuban was admitted into my wards on 16th February, 1916 He was cachectic and much emaciated at the time of admission The spleen extended 3" below the costal arch and there was a large number of L. D bodies present in the splenic blood. He was at first treated with intramuscular and subsequently with intravenous injections of colloidal metallic antimony. No unpleasant symptoms followed the intravenous injections. It was found that very quickly satisfactory results followed the treatment as will be seen from the following.

RESULT OF BLOOD EXAMINATION

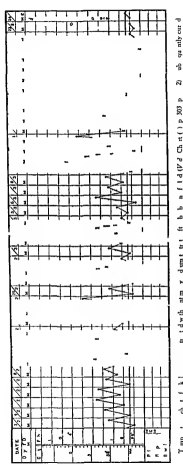
RE	3 C2,600,000	W E	3 C -4,800	Hb —40%	on	17-2-16
RE	3 C4,100,000	WE	3 C 5,600	Нь —50%	on	24-2-16
RE	3 C4,000,000	W E	3 C6,600	Нь —54%	on	6-3-16
RE	3 C-4,200,000	W E	3 C -7,800	Нь —54%	on	14-3-16
R I	3 C —4,300,000	W F	3 C 9,200	НЬ —70%	on	30-3-16

Size of spleen—3" below the costal margin in the left nipple line on 20-2-16 and 1'' on 14-3-16

Doses of colloidal antimony given -

Four intramuscular injections of 001 grm on successive days and 19 intravenous injections of 002 grm on the next successive days. The 20th injection was given eight days after the 19th injection.

The second case treated with colloidal antimony was Amiya She was at first treated with intravenous injections of berberine sulphate and subsequently with sodium antimonyl tartrate. As will be seen below, she got no benefit from berberine. The administration of sodium antimonyl tartrate was followed by severe diarrhea l, therefore, decided



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by c || d | m t || ankmony Patent-Am y - I t par



to put her on colloidal antimony and as will be seen below the results so far have been very satisfactory

Four Injections of Sodium Antimonyl Tartrate

Treatment with Colloidal Antimony

She had altogether 15 injections of 002 grm and 5 injections of 003 grm of the colloid intravenously

Observations

From the above it will be seen that the results so far seem to be very encouraging On one occasion the patient Bhuban was given 004 grm of the colloid and this was followed by a sharp rise of temperature The patient left hospital markedly improved in health and in body weight On the day of discharge the spleen could not be felt below the costal arch The second case is still in the hospital

- (b) Cases treated with metallic antimony in a state of fine subdivision as an impalpable powder —
- I have some more cases to report which were treated with this drug since the publication of my last paper. This would make a series of twelve cases which have been treated with metallic antimony alone or with metallic antimony after other drugs were tried.
- (1) Abdul —The patient has been free from fever for nearly four months He had altogether seven injections of metallic antimony once a week starting with ½ gr and ending with 13 grs [In my last paper it was reported that

treatment with metallic antimony was just begun in this case.]

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R B C -- 3,200,000
                   W B C - 4,000
                                                      5 11 15
                                               on
R B C -- 3,600,000
                   W B C - 6600
                                    Hb - 62\%
                                                     23-11 15
                                                on
                   W B C-10 200
R B C -4,700,000
                                    Hb -70%
                                                     14 12-16
                                               on
R B C -4,800 000
                   W B C-10,600
                                     НЬ
                                          82%
                                                      4 1 16
                                                on
                    W B C - 9.600
R B C -5,600,000
                                     Hb -76%
                                                      16 2-16
                                                on
```

Size of spleen -5'' below the costal margin on 21-10 15 and $\frac{1}{2}''$ on 16-2-16

(2) Abdul Aziz—was for some time treated with intravenous injections of narcotine. As the patient was not improving except in the leucocyte count and as the L D bodies were present even after ten injections of the drug, he was put on metallic antimony. He had altogether three injections of metallic antimony, each dose being one grain (Vide Temperature Chart.)

RESULT OF BLOOD EXAMINATION

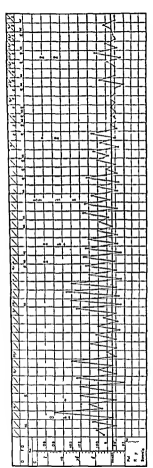
Treatment with Narcotine

R	B C2,900,000	W B C -2,400	Hb46%	on	13-1-16
R	B C3,400,000	W B C -4,600	Hb —48%	on	4 2-16

Treatment with Metallic Antimony

R B C -3,600,000	W B C — 5,400	Hb —50%	on	14-2 16
R B C -4,100,000	W B C — 8,200	I Ib -56%	on	23 2-16
R B C -4,500,000	W B C 10.400	Hb60%	on	4 4-16

(3) Patient—N.—He was at first treated with tartar emetic and his case was reported in the December number of the *Indian Medical Gazette* He was discharged from hospital much improved on 5th October, 1915, and was readmitted on 8th November, 1915 He had altogether four injections of metallic antimony, first two doses being one grain and last two doses one grain and a half each



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(4) Ascrit —He was at first treated with eusol. But as this was followed by slight improvement he was given 3 injections of metallic antimony. As a result of this the spleen which at first extended 5' below the costal arch could not be felt there.

RESULT OF BLOOD EXAMINATION

R B C - 4 200 000 W B C - 3 200 Hb - 58 / on 13 2 16 R B C - 4 900 000 W B C - 7 800 Hb - 66 % on 10 4 16 R B C - 4 900 000 W B C - 12 400 Hb - 66 % on 18 4 16

(5) Saha—He had four injections of metallic antimony. The blood count was RBC—2 600 000 WBC—2 400 Hb—35% before treatment and RBC—4 500 000 WBC—10 000 Hb—70% after treatment.

Observations

It was seen that every one of the above cases treated with intravenous injections of metallic antimony markedly improved under the treatment the improvement being noticed under the following heads (1) improvement of blood condition (2) diminution in the size of spleen (3) subsidence of the fever and (4) absence of the L D bodies from the spleen So far therefore these cases may be considered as cured In the above series of cases I have not taken into account a case which died of chronic dysentery five days after he had one injection of metallic antimony

1 Cases treated with Compounds of Antimony

A Cases treated with Soluble Salts of Antimony Tartar Emetic and Sodium Antimonal Tartrate

Since my earlier communications on the treatment of kala azar with antimony and its salts. I have treated some more cases with sodium antimony! tartrate either alone or combined with tartar emetic.

It was observed that most of these cases improved under the treatment, the improvement being noticed under the same heads as described above (see under I, pp 291-95) The total number of injections given to some of my cases are enumerated below —

- (1) M (already reported)—7 injections of sodium antimonyl tartrate (up to the last report) and 7 injections since
 - (2) Nandy—12 injections of sodium antimonyl tartrate
 - (3) Sing-20 injections of sodium antimonyl tartrate
- (4) Das (already reported)—7 injections of tartar emetic+10 injections of sodium antimonyl tartrate
- (5) Kamala (already reported)—6 injections of tartar emetic +8 injections of sodium antimonyl tartrate
 - (6) Satya—6 injections of tartar emetic
- (7) Kar—11 injections of tartar emetic+5 injections of sodium antimonyl tartrate
- (8) Saha—8 injections of sodium antimonyl tartrate (treatment discontinued and patient put on metallic antimony, as there was no improvement in the blood condition after 8 injections)
- (9) Sishir—19 injections of sodium antimonyl tartrate (treatment still continued)
- (10) Souren—9 injections of sodium antimonyl tartrate + 2 injections of tartar emetic (treatment still continued)

One of my youngest cases treated with sodium antimonyl tartrate was a patient aged 3 years. Up to now, the highest dose she has received is 2 c c of 2 per cent solution of the salt. She is now being treated with the sodium and the potassium salts alternately and is progressing favourably

The injections were stopped after the patient had been free from fever for some time, the spleen had gone down to almost under the costal arch, the Leishman-Donovan bodies had disappeared from the splenic blood and the leucocyte count had been high

The results of examination of blood in some of my cases have been shown in my previous papers. All the cases that have been already reported in the papers have since been free from fever and have markedly improved in health. The latest report of examination of blood of some of these cases treated with the soluble salts is as follows.—

- (1) Patient M (reported in the January number of the Indian Medical Gazette), R B C -4 500 000 W B C -8 500 Hb -75% on 15th February 1916 (patient had altogether 14 injections of sodium antimonyl tartrate)
- (2) Das (reported in the above paper) R B C 4 500 000 W B C 9 000 Hb 75% on 16th February 1916 (patient had 7 injections of tartar emetic and 10 of sodium antimonyl tartrate)
- (3) Nandy—R B C —2 800 000 W B C —4 600 11b —38% on 20th December 1915 R B C —5 200 000 W B C —8 500 Hb —70% on 28th February 1916 (patient had 12 injections of sodium antimonyl tartrate)

Observations

lt requires to be seen whether any of the cases mentioned above would get a relapse after some months. If they do not then they may be considered as cured. The doses and the intervals between successive injections were described in my previous papers. The alternate administration of the sodium and the potassium salts seems to be attended with the best results.

Metallic Antimony and its Soluble Salts compared

lt is at present difficult to state from chinical experience whether metallic antimony or its soluble salts would constitute the best form of treatment in hala azar though as stated in my previous papers from theoretical grounds and from comparison with trypanosomiasis, metallic antimony holds

out the best chances of cure The following facts have, however, at present, been observed —

- (1) The number of injections of metallic antimony required is much less than that of its soluble salts to produce the beneficial effects in kala-azar (Three to four injections are generally required)
- (2) The effect is more quick after injections of metallic antimony
- (3) The toxic symptoms were less marked and the beneficial effects more permanent after injections of metallic antimony than after injections of its soluble salts in the following case—

The patient Narayan (already reported in the December issue of the Indian Medical Gazette) was at first treated with tartar emetic and was apparently cured after seven injections He was discharged from the hospital on the 5th October, 1915. and was re-admitted on the 8th November, 1915 He still had an enlarged spleen, and though he was free from fever, his general condition seemed to have somewhat deteriorated He was at first given an injection of galyl which apparently did him no good Then he was given 3 c c of 2 per cent solution of sodium antimonyl tartrate This was followed by severe enteritis and could not be continued He was then put on a course of treatment with intravenous injections of metallic antimony, and this was quickly followed by remarkable improvement in the blood count and in general health The following is a short account of his blood count after the various treatments ---

R B C --2,800,000, W B C --1,600, Hb.--46% on 4-9-15

R B C —3,800,000, W B C —6,400, Hb —70% on 2-10-15 after 7 injections of tartar emetic

R. B. C.—3,600,000, W. B. C.—5,600 Hb.—62% on 11-11-15 (on re-admission into hospital).

R B C -3 400 000 W B C -4 400, Hb -58% on 21 11 15 (3 gramme galyl injected on 17 11 15)

R B C -3 300 000 W B C -6 400 Hb -54% on 20 12 15 (3 c c of 2 per cent solution of sodium antimonyl tartrate injected on 27 11 15)

R B C —3 800 000 W B C —13 800 Hb —68% on 17 1 16 (one grain of metallic antimony injected on 15 12 15 and one and a half grains on 26 12 15 and 4 1 16)

It will be seen in the above case that the greatest benefit followed the injection of metallic antimony

The comparison of the effects of colloidal metallic antimony with those of metallic antimony in a state of fine subdivision as an impalpable powder is reserved for a future communication.

Observations

The following facts may be mentioned here -

- (1) Colloidal metallic antimony is administered in very small doses (002 gramme)
- (2) The colloid may be injected in 2 c c suspension and is therefore most convenient to use
 - (3) No unpleasant symptoms follow the injection of colloidal antimony

B Case treated with Colloidal Solution of Sulphide of Antimony

One case is being treated with colloidal sulphide of antimony given subcutaneously in doses of 002 gramme

III TREATMENT WITH INTRAVENOUS INJECTIONS OF ANTISEPTICS

(a) I have already reported an apparently cured case after treatment with intravenous injections of formalde hyde. Since the above case was published two more cases have been treated with the same drug

(1) The patient, named Kası, is one of them and was reported before. As will be seen from my previous paper, though the patient got rid of his fever and his general condition improved after two injections of metallic antimony, his blood count was still below the normal I decided to try intravenous injections of formaldehyde and a remarkable improvement in his blood followed, as will be shown below —

```
R B C -2,600,000, W B C -5,000, Hb -55% on 28 11-15
R B C -2,600,000, W B C -4,200, Hb -50% on 3-12-15
R B C -3,700,000, W B C -9,800, Hb -62% on 4 1 16
```

Doses of Formaldehyde given

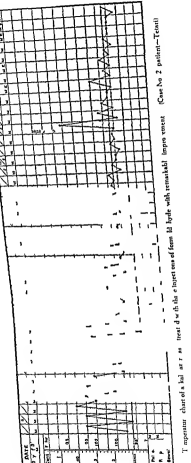
18 c c	of	4000	solution	of formaldehyde	injected	on	5 12-15
36 с с	of	do	,,	*1	,,	on	6 12 15
35 с с	of	3000	,,	11)1	on	7-12 15
36 с с	of	do	,,	1)	,,	on	9-12-15
34 c c	of	2000	,,	,,	,,	on	14-12-15

(2) The patient, named Tetari, was admitted on 17-1-16 into my wards. She was given three injections of formal-dehyde and as a result of this there was a remarkable improvement in her condition. (Vide Temperature Chart.)

```
R B C -- 2,500 000, W B C -- 3,000, Hb -- 42% on 24-1-16
                    W B C - 3,800, Hb -42% on 31-1-16
R B C -- 2,600 000,
R B C -4,000,000.
                   W B C - 11,200,
                                      Hb -54% on 7-2 16
R B C -3,900,000,
                    W B C - 7,900,
                                      Hb -56% on 15-2-16
Size
      \mathbf{of}
          spleen—4"
                       below the costal
                                        margin on 24-1-16
Size
      \mathbf{of}
          spleen—3//
                       below the costal margin on 20-2-16
```

Doses of Formaldehyde given

60 c c	of	4000	sol d	of formaldehyde	ınjected	on 27-1-16
50 с с	of	3000	,,	,,	,	on 28 1-16
32 c c	of	2000	,,	,	,,	on 31-1-16
35 сс	of	<u> 2000</u>	,,	,,	,,	on 16-2-16





(b) The next antiseptic used was eusol prepared afte-Lorrain Smith's formula

The patient named Ascrit was admitted into my wards on 17th January 1916. The spleen extended 5' below the costal margin in the parasternal line and a large number of L. D. bodies were found on spleen puncture. He was given altogether 15 injections of eusol. There was a slight improvement in the blood count but the results were not so satisfactory as the e following treatment with antimony as will be seen from the following notes.—

```
R B C - 3 500 000 W B C - 3 800 Hb - 48, on 19 | 16
R B C - 3 900 000 W B C - 4 200 Hb - 50 / on 3 | 1 | 16
R B C - 4 200 000 W B C - 5 800 Hb - 58 / on 19 2 | 16
R B C - 4 300 000 W B C - 4 800 Hb - 60 / on 2 3 | 16
Size of spleen-not duninghed
```

Doses of Eusol given

The doses varied from 55 cc to 200 cc of eusol given intra venously on successive days

As stated above the patient was subsequently treated with metallic antimony with brilliant results

It will be seen that formaldehyde acted like a specific in the three cases in which it was used the effect being noticed under the same heads as those described under antimony and its salts. Evidently however the drug must be given a more extended trial before its specific action is established beyond doubt. In the single case mentioned above in which cusol has been tried the results so far seem to be slightly promising but much less marked than with formaldehyde lt may be mentioned here that there were no untoward results following intravenous injection of cusol. The objections to formaldehyde however he in the fact that it is liable to decomposition and different samples contain different strengths. It was therefore decided to use a preparation of tormaldehyde which is more stable and obtainable in

crystalline form, and this is formaldehyde-sodium bisulphite. A case is being treated with this drug and will be reported later on

IV ALKALOIDAL THERAPY

Attempts were made to discover an alkaloid which would exert a specific action in kala-azar

In one case, quinine was given intravenously in the form of quinine bihydrochloride but no impression was made on the course of the disease

In another case, already mentioned, Abdul Aziz, narcotine was used intravenously. The effect on the blood was to some extent beneficial, as will be shown by the tables given under I, but there was no effect on the temperature. The patient was afterwards put on a course of intravenous injections of metallic antimony. Another case has been treated with the drug and so far the results are the same as in the above case.

It may be noted here that it is somewhat difficult to put narcotine into the veins as it is insoluble in water as well as in normal saline

Doses of narcotine —(1) ½gr given intravenously on 13-1-16, (2) ½gr on 14 | 16, (3) ½gr on 15-1-16, (4) | gr on 17-1-16, (5) | gr on 19-1-16, (6) | gr on 21-1-16, (7) | gr on 23-1-16, (8) | gr on 26-1-16, (9) | ½grs on 3 | 1-16, (10) | ½grs on 2-2-16

As just now stated, the effect of treatment with narcotine was only an improvement in the blood condition but there was no effect on the temperature and the L D bodies were present in the spleen

The next alkaloid used was berberine sulphate I have already reported that the combination of this drug with antimony does not seem to lead to better results than what

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nperature chart of a kala azar case, treated with berberine sulph intravenously but discontinued 18 there was no improvement (Para 2, patient—

is obtained from antimony itself (Indian Medical Gazette January 1916) Two cases were treated with intravenous injections of this drug. One of them died of pneumonia and on post mortem examination L. D. bodies were found in the spleen. He was given five injections of berbenne sulph intravenously the dose being ½ to 13 grains.

The next patient named Amiya was at first treated with intravenous injections of berberine sulph. She was given altogether 10 injections. There was no improvement in the blood count and there was no diminution in the size of the spleen and a very large number of L. D. bodies were present in the splenic blood after these injections. As the patient was not improving and was getting fever berberine sulph was discontinued. The drug was given in 1 grain doses for seven days and in 2 grain for three days.

Result of Blood Examination

R B C -- 4 300 000 W B C -- 2 400 Hb -- 44% before treatment

R B C -4 000 000 W B C -3 200 Hb -46% after treatment with berberine sulphate

The next alkaloid that has been used is bebeerine sulph. The effect of treatment with this drug has been unsatisfactory.

I have observed the following unpleasant effects after injections of sodium antimonyl tartrate or tartar emetic in some of my ca es (1) Rigor and temporary rise of temperature (2) pain in the gums (3) vomiting (4) a troublesome cough followed by vomiting in some cases (5) in one case there was an intense headache which lasted for nearly three days after injections of sodium antimonyl tartrate (6) diarrhoca

Conclusions

(1) Colloidal metallic antimony has been used intrave nously for the first time in kala ezar with very promising

results in the cases in which it was used It is administered in very small doses.

- (2) Antimony, in a state of fine subdivision, has been given intravenously with brilliant results in kala-azar
- (3) Tartar emetic and sodium antimonyl tartrate have also been used with marked benefit in the disease. The alternate administration of the two salts seems to be attended with the best results
- (4) The number of injections of metallic antimony required for a course of treatment is very much less than that of the soluble salts.
- (5) Formaldehyde has been used in three cases with excellent results. The effects of eusol are much less marked than those of formaldehyde
- (6) Intravenous injections of narcotine is followed by an improvement in the condition of the blood Intravenous injections of berberine sulph and bebeerine sulph, have not given rise to any beneficial results in the cases in which they were tried
- (7) Antimony or its combination with formaldehyde will perhaps constitute a treatment in kala-azar and will cure a large percentage of cases of this fell disease which is deadlier than tuberculosis and kills hundreds of people in Assam and many parts of Bengal
- (8) If future observations confirm the view that three or four injections of metallic antimony are sufficient to bring about a complete and permanent cure of the disease, then we are in possession of a drug as powerful as quinine is for malaria, emetine for amoebic dysentery or salvarsan for syphilis. Its combination with formaldehyde will, perhaps, still more cut short the duration of the disease by the destruction of any antimony-fast parasites that may come into existence.

FOURTH REPORT ON THE TREATMENT OF KALA AZAR AND SOME BLOOD REACTIONS IN THIS DISEASE

Α

FOURTH REPORT ON THE TREATMENT OF KALA AZAR

Cases of enlarged spleen admitted into my wards from July to October in one year and whose splenic blood I examined numbered forty. Out of these twelve showed the parasite of kala azar in the splenic blood. These together with a series of cases published by me in the British Medical Journal in May 1908 show that out of one hundred and ninety cases of enlarged spleen in which the splenic blood was examined the L. D. bodies were found in 72 of them or 38 per cent.

In another series of one hundred and sixty six cases the results of examination of the splenic blood were as follows —

- (1) 27 1 per cent showed L D bodies
- (2) 21 1 per cent showed malarial parasites
- (3) 012 per cent showed both malarial parasites and L. D. bodies
 - (4) 51 8 per cent showed neither malarial parasites nor

L D bodies

Taking all the three series into account it will be found that out of three hundred and fifty six cases in which the splenic blood was examined by me only one hundred and eighteen or 33 per cent had the parasite of kala azar

The extreme rarity of malarial parasites being found with L D bodies is noteworthy and points to the conclusion that the two diseases rarely go together

The results are also striking, in view of the fact that malaria and kala-azar cannot account for the enlargement of the spleen in a large percentage of the cases met with in the medical wards in Calcutta

The cases with negative results in the examination of the splenic blood sometimes, though rarely, gave a blood picture resembling that of kala-azai. The notes of one such case are appended below —

Patient, æt 13, was admitted with the spleen extending $9\frac{1}{2}$ inches below the costal margin and with a history of fever for about two years. The spleen was punctured three times but neither any malarial parasites nor any L. D. bodies were discovered after very careful examination. The blood count was as follows—

(1) R. B C	2,260,000
(2) W. B. C.	1,500
(3) Hb.	40 per cent
(4) $\frac{W. B. C.}{R B C} = \frac{1}{1.540}$	

Differential count —

Polymorphonuclears	47 per cent
Large mononuclears	13 ,,
Lymphocytes	36 ,,
Eosmophiles	4

The temperature was normal throughout his stay in hospital, except on two occasions when the temperature rose to 103°F. The blood was examined on these occasions but neither malarial parasites nor L. D. bodies were discovered. The patient was in hospital for more than three months. Cases like the above are not infrequently met with.

From the above statistics one is led to think of the possibility of an undiscovered though common cause of splenic enlargement in India. Are some of them cases of splenic anæmia or is it possible that there is an yet undiscovered phase of leishmania in which they disappear from the spleen and reside in some other parts of the body and in this quiescent stage give rise to httle or no febrile manifestations or are they due to some undiscovered parasites?

The treatment of kala azar described in this paper can be divided under the following heads —

- (1) Treatment with tartar emetic and sodium antimonyl tartrate given intramuscularly
- (2) Treatment with lithium antimonyl tarirate given intramuscularly
- (3) Treatment with aniline antimonyl tartrate given intravenously
- (4) Treatment with antimonious oxide in a state of fine subdivision given intravenously
 - (5) Treatment with luargol
 - (6) Treatment with colloidal oxide of antimony
- (7) Treatment with intravenous injections of bismuth tartrate solubilis

Treatments under (1) to (5) have been described in detail in my treatise Kala azar Its Treatment published by Butterworth & Co (India) Ltd They will not therefore be described in detail here The following conclusions can be drawn from them —

- (1) Cases treated with intramuscular injections of tartar emetic and sodium antimonyl tartrate improved remarkably. The injections were painful (dose 1 to 11 c c of 2 per cent solution given every 4th or 5th day)
- (2) Cases treated with lithium antimonyl tartrate given intramuscularly showed much improvement in blood condition but patients left hospital before treatment was completed (dose 1 to 3 gramme) The injections were painful

- (3) Cases treated with aniline antimonyl tartrate, given intravenously, recovered completely (dose 1 to 10 c c of 2 per cent solution) The febrile reactions as well as vomiting, purging or rigors were much less common with this salt than with potassium or sodium antimonyl tartrate I cannot, at present, state whether it is superior to sodium or potassium antimonyl tartrate in its therapeutic effects
- (4) Cases treated with intravenous injections of antimonious oxide in a state of fine subdivision were subsequently treated with metallic antimony and recovered completely (dose—75 gr to 1.5 grs)
- (5) Cases treated with luargol showed improvement in the blood condition, but the parasites persisted. Patients left hospital before treatment was completed
 - (6) Cases treated with colloidal oxide of antimony

In the Addendum to my Treatise on Kala-azar a reference has been made by me to the use of colloidal oxide of antimony. It seems to be the least toxic of all the antimonial preparations that I have used in kala-azar. It is at present difficult to compare its therapeutic value with that of other antimonial preparations. The following are the notes of one case treated with this drug

Patient A—was admitted into my ward with the spleen extending $2\frac{1}{2}$ inches below the costal arch. He was treated with intravenous injections of colloidal oxide of antimony, beginning with 4 c.c of the colloid and increasing the dose by 1 c c at successive injections. For some time the injections were given every day. The patient has up to now had altogether eleven injections, the highest dose being 10 c c. The effects of the treatment were as follows:—

Body weight-same.

Blood ·—

R B C -2,600,000, W B C -1,200, Hb -40% on 19-4-17 before treatment

RBC —3 000 000 WBC —2 400 Hb —38% on 2 5 17 during treatment

RBC-3 300,000 WBC-4 000 Hb-48° on 12 6 17 during treatment

There seems to be a stendy but slow improvement in the general condition of the patient ond the drug seems to be perfectly free from any toxic symptoms

The high temperature which the patient had before treatment come down there being only a slight rise towards the evening lasting for a very short period

(7) Cases treated with intravenous injections of bismuth tartrate solubilis (bismuth sodium tartrate)

In o paper rend before the Royal Society of Medicine 9th January 1909 Cushny pointed out that arsenic anti-mony ond bismuth killed trypanosomes in the concentration of 1 200 000 According to his observations bismuth proved too poisonous to the hest and the damage done was irreparable

I have given bismuth tartrate solubilis intravenously in a few cases of kala azar. The toxic symptoms observed by Cushny in experimental trypanosomiasis were not noticed in any of my cases of kola azar and in one case there was such a remarkable improvement in the patient's condition that there was every chance of his perfect recovery. The preliminary notes of this case have been published in my Treatise on Kala azar already referred to I append here the full notes of this case giving further observations since the publication of the above work.

Patient B—was admitted into my ward on the 4th February 1917. The spleen extended 6 inches below the costal arch and L. D. bodies were found on spleen puncture. The patient was emaciated and cachectic. He was at first treated with one per cent solution of bismuth tartrate solubils in distilled water beginning with 1 c.c. and increasing up to 10 c.c. He was then treated with 5 c.c. of 2 per cent

solution, the dose being gradually increased to 9 c c. Altogether 17 injections were given. As a result of this treatment there had been a remarkable improvement in the condition of the patient as shown by the following note:—

- (1) Increase in body weight—22 lbs in two months, during which the treatment was continued.
 - (2) Size of the spleen—diminished by 4 inches
 - (3) Disappearance of the fever
- (4) Marked improvement in the blood condition as shown by the following figures —

It will be seen that there was at first a diminution of the leucocytes during the treatment, but afterwards they steadily increased The patient is now enjoying perfect health and has recently returned to work.

Two more cases were treated with the same drug, but they left hospital before treatment was completed. In one of these, the patient complained of intense pain in the gums each time the injections were given and there was marked pigmentation in the gums. The symptoms that are frequently met with after injections of antimonial preparations such as vomiting, cough, rigor, high fever and sudden syncope were completely absent in all these cases.

I hope to give at a future date a statement showing the comparative value of antimony and bismuth in the treatment of kala-azar

R

A PRELIMINARY REPORT ON SOVIE BLOOD REACTIONS IN LALA AZAR

l The relative hæmoglobin value of the resistant

In the Bio Chemical Journal Vol 1V 1909 1 described a new method of testing blood in a paper entitled. The relative haimoglobin value of the resistant erythrocytes during hæmolysis of blood etc. I have subsequently found that this factor is markedly diminished in kala azar as will be seen from the following figures —

		I part of blood +2 part I di tilled water
(I) F	lealthy student	372
(2)	Do	434
(3)	D _o	428
(4)	Do	450
(5)	D _o	448
		Average = 426
(1) 1	lalarial fever	606
(2)	Do	303
(3)	Do	444
		Average = 451
(1) 4	Ankylostomiasis	714
(2)	Do	658
		Average = 686
(1)	Kala azar	266
(2)	Do	250
(3)	Do	285
(4)	Do	142
		Average = 236

The diagnostic importance of these facts can be settled by further observations

II The complement deviation reaction in kala azar—

A limited number of observations have been made by me in this direction. In a series of eight cases, the reaction was

found to be positive in six and negative in two. The diagnosis of each case was made by spleen puncture and the antigen used was made by the alcoholic extract of the fresh spleen of a kala-azar case made by grinding up one part of the spleen pulp with three parts of a mixture of equal parts of alcohol and '85 per cent NaCl solution and then heated for an hour at 60°C. As the test originally employed by me was Fleming's modification of Wassermann test, and as Fleming's test is not free from fallacies, I do not consider any importance can be attached to these results till further observations have been made by more accurate methods. So far, these results differ from the observations of Pavoni in infantile kala-azar.

Very recently I tested this reaction according to the original Wassermann method and found that in only one out of four kala-azar cases this reaction was positive

III Hæm-alkalınıty — Archibald was the first to point out that the alkalınıty of blood is diminished in kala-azar. Rogers and Shorten have confirmed this in Indian kala-azar.

I have tested the blood in a series of kala-azar cases for determining the basic reactivity. This was determined by a modification of that described by Moore and Wilson (Bio-Chemical Jouinal, 1909) Fifty c.mm of blood were taken from the finger which was sterilized by 5 per cent formol solution and put into a perfectly dry sterilized tube and then quickly centrifuged Ten c.mm of the serum were treated with a solution of $\frac{N}{100}$ H₂SO₄ in a white porcelain shallow capsule, using a drop or two of a fresh dilution of an alcoholic solution of dimethyl-amido-azobenzol in distilled water, the first indication of neutralization being given by a faint rose colour at the side of the liquid in the porcelain capsule. The dimethyl is first dissolved in alcohol and then one drop of this is mixed with distilled water just at the time of the experiment drops of this are added to the serum to make it faint yellow.

In a series of cases the average basic reactivity was found to be 092 normal as compared with 178 normal in a series of healthy students (see my Treatise on Studies in Hæmolysis Calcutta University Series)

IV $Ham \ salinity$ —This was estimated by treating 10 c mm of the serum with $\frac{N}{2}$ AgNO₃ using a solution of K CrO₄ as an indicator. The average was found to be 6538 per cent as contrasted with 6654 per cent in a series of observations on healthy students

V It is frequently observed that when the blood of a kala azar patient is mixed with excess of distilled water a white flocculent precipitate forms. While this reaction is present in a large majority of cases of kala azar it has also sometimes been observed in other diseases of phthisis, cancer of the liver cirrhosis of the liver chronic malaria and cases of enlarged spleen in which no L. D. bodies were found on spleen puncture

It has also been found that this precipitate is obtained by mixing the serum separated from the corpuscles with excess of distilled water. The red corpuscles when dissolved in distilled water do not give rise to any such precipitate precipitate is soluble in a solution of sodium bicarbonate, and also in dilute acetic acid as well as in normal saline insoluble in distilled water. It seems that the precipitate is of the nature of a globulin and is probably easily precipitated on account of the diminished alkalimity of the blood in kala azar Whether it is due to the presence of any excess of globulin in the serum of kala azar cases due to dis integration of leucocytes or whether this globulin is of a specific nature has not yet been determined. As regards its diagnostic importance it may not be of great value because as stated before a similar precipitate has been found in other diseases

One remarkable property of this globulin-like substance is that a solution of it in normal saline inhibits the action of the complement in a hæmolytic system

My grateful thanks are due to Lt. Col W D Sutherland, I.M.S., Imperial Serologist, and his assistant Dr G C Mitter for providing me with materials for my hæmolysis work

ON THE PRESENCE OF AN EASILY PRECIPITABLE ANTI-COMPLEMENTARY GLOBULIN LIKE SUBSTANCE IN HUMAN SERUM AND ITS IMPORTANCE IN THE DIAG NOSIS OF KALAAZAR

When human serum is diluted with excess of distilled water it becomes cloudy owing to partial precipitation of the serum globulin. Under certain circumstances a copious precipitate forms instead of mere cloudiness. This precipitate is due to a globulin like substance as is evident from the following facts—

- (1) It is soluble in normal saline in dilute acids and in dilute sodium bicarbonate solution — It is also soluble in sodium hydrate solution
- (2) It is precipitated from its solution in normal saline when the solution is treated with equal parts of a saturated solution of (NH₄) SO₄ or when it is saturated with MgSO₄ or NaCl
- (3) It is not precipitated by NH₄OH from its solution in dilute acids
 - (4) It is insoluble in distilled water

On chemical analysis, this substance is found to contain C, N, H, O, but so far I have not been able to detect in it the presence of S, P, or any halogen. After being thoroughly washed in distilled water it can be collected as a white precipitate having a granular appearance under the microscope.

If further investigations confirm the observation that this substance does not contain any phosphorus or sulphur, then it will be found to be different from serum-globulins in chemical composition.

I hope to enter, at a future date, into the chemical nature of this substance and at present shall content myself by assuming that it is globulin-like in nature

In the Indian Medical Gazette, September, 1917, I pointed out that this copious precipitate is frequently observed when the serum of a kala-azar case is mixed with excess of distilled I also pointed out that a precipitate apparently similar to this has sometimes been observed in other diseases, e.g., chronic malaria, phthisis, cancer of the liver, etc I was not, therefore, then able to state whether the presence of this precipitate was of any diagnostic importance ın kala-azar Further observations lead to the conclusion that, if instead of using an excess of distilled water (which in my original experiments consisted of 15 to 20 times the amount of serum used) one uses two or three volumes of distilled water then the precipitate appears almost exclusively ın kala-azar Thus in a series of 20 cases of kala-azar, the following results were obtained —

One part of serum plus two parts of distilled water produced a copious precipitate. In some cases one part of serum plus one and-a-half parts of distilled water gave rise to a distinct precipitate

Similar experiments were made with the serum of a series of cases suffering from other diseases and a negative result was always obtained

On p t of um plu tw p t of d t ll d w t

No pp

(1)	Phthisis	No pp
(2)	Malarial fever	No pp
(3)	Cirrhosis of the liver	No pp
(4)	Enteric fever	No pp
(5)	Bright's disease	No pp
(6)	Ankylostomiasi	No pp
(7)	Pernicious anaemia	No pp
(8)	Dengue	No pp
(9)	Dysentery	No pp
(10)	Pneumonia	No pp
(11)	Catarrhal jaundice	No pp
(12)	Broncho pneumonia with enlarged	
	spleen (no L D bodies in the	

In a few cases with enlarged spleen in which no L. D bodies were found on spleen puncture a similar precipitate was obtained though clinically they looked like kala azar

spleen)

Whether these are cases of kala azar in which the parasites could not be found on spleen puncture as some times is the case as pointed out by Leishman or whether some of them are cases of spontaneous cure from kala azar cannot be definitely stated in the present state of our knowledge

I have also found that if distilled water is gently poured on the top of the serum of a kala azar case a distinct white ring is formed at the junction similar to the ring of albumin that is found on addition of nitric acid to a solution of albumin. This test also appears to be of diagnostic importance in kala azar. A similar ring is also observed in some obscure cases of enlarged spleen mentioned before

To perform the above two tests proceed as follows -

(1) Two c c of the blood from a prominent vein of a kala azar case are drawn by a glass syringe and the blood quickly centrifuged. The serum freed from the clot is introduced into a miniature test tube with a capillary pipette and then a small amount of distilled water is gently poured over the serum. A distinct white ring forms at the junction in every case of kala-azar. I propose to call this the "globulin ring test" of kala-azar.

(2) The serum is collected in a miniature test tube and then mixed with two or three parts of distilled water. A white precipitate forms in every case of kala-azar. I propose to call this the "globulin precipitation test" of kala-azar.

The Anticomplementary Properties of the Globulin-like Substance

This fact was briefly touched upon by me in the September number of the *Indian Medical Gazette* The following is a detailed method of showing this remarkable property of this globulin-like substance —

One c.c of the serum of a kala-azar case is mixed with fifteen c c. of distilled water. The precipitate is collected, washed thoroughly with distilled water and then dissolved in one c.c. of normal saline. The following observations were made:—

- (1) Take, for instance, a hæmolytic system in which the following are the doses of the component parts—
- '15 c.c. of anti-sheep amboceptor + 5 c c of guinea-pig's complement + '5 c.c of sheep's corpuscles = complete hæmolysis
- (2) (a) Mix 5 c c of guinea-pig's complement with 2 c c of the solution of the above precipitate in normal saline and incubate for half an hour.
- (b) Add to this 15 c c of anti-sheep amboceptor + 5 c c: of sheep's erythrocytes = no hæmolysis (I propose to call this test the "anticomplementary globulin test" of kala-azar)
- (3) 1 c c of the serum of the same kala-azar case + 5 c c. of sheep's erythrocytes = complete hæmolysis (due, no

doubt to the natural complement and amboceptor frequently present in human serum)

- (4) (a) Heat the serum to 55°C for half an hour
- (b) Add 2 c c of the heated serum to 5 c c of guinea pig complement and incubate for half an hour
- (c) Add 5 c c of sheep s erythrocytes to (b) and incubate = complete hæmolysis

From the above the following conclusions are drawn -

- l One part of serum of a kala azar case with two or three parts of distilled water gives a distinct precipitate. Such a precipitate is not obtained in any other disease except kala azar and some rare obscure cases of enlarged spleen. Its presence is therefore of much diagnostic importance (globulin precipitation test)
- 2 Gently pour distilled water on to the top of the serum of a kala azar case a distinct white ring forms at the junction. This ring is not observed in any other disease except kala azar and some rare obscure cases of enlarged spleen. Its presence is therefore of much diagnostic importance (globulin ring test)
- 3 The solution in normal saline of the above globulin like substance present in the serum of kala azar patients inhibits the action of the complement in a hæmolytic system consisting of sheep's corpuscles anti-sheep amboceptor and guinea pig s complement (anticomplementary globulin test)
- 4 This globulin like substance does not inhibit the action of the natural complement normally present in the serum as long as it is not separated from the serum by the action of distilled water
- 5 This globulin like substance is probably in combination with some constituents of the serum and as long as this combination exists it exerts no anticomplementary action Distilled water also breaks up this combination. It is not broken up by heating the serum to 55°C.

- 6 We have regarded the above substance to be globulinlike in nature, but further investigations will be required to determine its chemical nature. At present it seems to differ from serum-globulin in not containing S or P.
- Globulin-like substances are sometimes precipitated from the serum of cases suffering from chronic malaria, cancer of the liver, etc., by the addition of excess (15 or 20 parts) of distilled water, as has been pointed out by me in the Indian Medical Gazette, September, 1917 The precipitate obtained is much greater than, and must not be confounded with, what appears as a cloudiness when normal serum is diluted with excess of distilled water, due to partial precipitation of serum globulin None of these, however, are precipitated when the serum is diluted with only two or three parts of distilled water. The properties of these globulin-like substances will form the subject of a future It cannot, at present, be stated whether they investigation possess any anticomplementary properties similar to what is shown by the globulin-like protein separated from the serum of kala-azar patients I cannot also state whether they contain any P or S
- 8 It is possible that in different diseases, globulinlike proteins, perhaps of a specific nature, are present in the serum with varying degrees of solubility in salt solution. The one present in kala-azar is characterised by being very easily precipitated by addition of a small amount of distilled water to the serum. This is evidently due to the inability of NaCl in the diluted serum to hold the protein in solution.

Whether the anticomplementary globulin test, the globulin precipitation test and the globulin ring test are absolutely pathognomonic of kala-azar can only be settled by further investigations, but, so far, they seem to be very valuable tests in the diagnosis of the disease. In some cases of very slightly enlarged spleen, the production

of these reactions led to the diagnosis of kala azar which was afterwards confirmed by spleen puncture ln making the ring test the serum must first be diluted ten to twenty times with normal saline

My grateful thanks are due to Lt Col R P Wilson I M S for giving me every facility in carrying on my researches in the Campbell Hospital I am also deeply indebted to Col W D Sutherland I M S and his assistant Dr G C Mitter for providing me with materials for conducting the serological portion of this investigation

TREATMENT OF KALA-AZAR WITH INTRAMUSCULAR INJECTIONS OF HYPER-ACID ANTIMONYL TARTRATE (+ URETHANE)

Since the discovery of antimony as a specific in the treatment of kala-azar, attempts have been made to discover a preparation which could be given intramuscularly without local reaction. The ordinary antimonyl preparations, such as tartar emetic or sodium antimonyl tartrate, give rise to violent local reaction and cannot therefore be used intramuscularly.

Caronia has used acetyl-p-aminophenyl-stibinate of sodium intramuscularly in the treatment of infantile kala-azar with good results and subsequently it was used by Kharina-Marinucci

In seeking for a preparation of antimony which will give little local irritation, we should use one which will be quickly absorbed without dissociation or decomposition. Such a preparation I have found in hyper-acid antimonyl tartrate (+ urethane). It is readily soluble in water, stable in aqueous solution for indefinite periods, and is quickly absorbed without decomposition after intramuscular injection. As urethane is not a base, it probably remains in solution with the antimonyl compound in the form of a mixture.

Experiments are being conducted by me to determine its toxic dose as compared with its curative dose, and, so far as I have been able to determine, it appears to be the least toxic of all the antimonial preparations and its curative

١,

dose seems to be much smaller than that of other antimonial preparations Further observations on this subject will be communicated in a future paper

The following are the series of the first four successive cases which have been treated successfully with this compound. In each of these cases the diagnosis was made by the presence of L. D. bodies in the spleen and the cure was shown by their disappearance therefrom —

1 Patient B S—was admitted into my ward on 25 9 19 with the spleen extending 6 in below the costal margin in the left nipple line. He was given intramuscularly 2½ c c of a two per cent solution of the hyper salt with urethane Altogether 14 injections were given from twice to four times a week. The results of treatment were as follows—

R B C —2 800 000 W B C —1 800 Hb —46 per cent on 26 9 19 before treatment

R B C —4 700 000 W B C —13 800, Hb —60 per cent on 5 I 20 after treatment

There was a marked increase in weight the spleen could not be felt below the costal arch no L D bodies could be found on spleen puncture and the fever subsided

2 Patient M —was admitted into my ward on 23 8 19 the spleen extending 5 in below the costal margin in the left nipple line. He was given 2½ to 5 c c of 2 per cent solution of the hyper salt with urethane intramuscularly Altogether 15 injections were given from twice to four times a week. The results of treatment were as follows —

R B C —3 300 000 W B C —2 200 Hb —38 per cent on 8 9 19 before treatment

R B C -4 600 000 W B C -16 000 Hb -60 per cent on 23 12 19 after treatment

There was a marked increase in weight the spleen could just be felt below the costal margin no L D bodies could be found on spleen puncture and the fever subsided

3 Patient R. B.—was admitted into my ward on 27-10-19, the spleen extending 3 in below the costal arch in the left nipple line. He was given 17 injections intramuscularly in doses of $2\frac{1}{2}$ c c of 2 per cent solution, every two to three days. The results of treatment were as follows —

R B C —3,900,000, W B C —2,200, Hb —46 per cent on 29-10-19 before treatment

R B.C —4,900,000, W B C —10,400, Hb —60 per cent on 19-1-20 after treatment.

There was marked increase in weight, the spleen could not be felt below the costal arch, the fever subsided and no L D bodies could be found on spleen puncture.

4 Patient B—was admitted into my ward on 6-11-19, the spleen extending 3½ in below the costal margin in the left nipple line. He was given only 5 injections of the hyper-salt with urethane at intervals of 3 to 4 days in doses of 2½ c c of 2 per cent solution.

The results of treatment were as follows -

R B.C.—3,100,000, W B C —2,400, Hb —48 per cent on 12-11-19 before treatment

R.B.C.—4,800,000, W B C —12,600, Hb —60 per cent on 20-1-20 after treatment.

There was marked increase in weight, the spleen could not be felt below the costal arch, no L D bodies could be found on spleen puncture and the fever subsided

As regards local irritation, there is, in some cases, some amount of swelling at the site of injections which subsides quickly. No abscess or necrosis was found in any of the cases. The highest dose was 5 c c of 2 per cent solution calculated in terms of the amount of Sb₂O₈ present. No reaction in the form of rigors, high fever, or cough was observed in any of the cases. The number of injections given, up to now, to all my cases was nearly 100

Another series of cases is being treated with the same compound

I have subsequently found that one per cent solution is almost absolutely painless. It appears to me that the use of the hyper acid antimonyl tartrate is a great advance in the treatment of kala azar

Very recently I have prepared urea and antimonyl tartrate and a trial is being given to it by intramuscular injections in kala azar. The results of these observations will be published in a future communication. So far it seems to be promising

A PRELIMINARY NOTE ON THE GLOBU-LIN, ALBUMIN AND CHOLESTEROL CONTENTS OF THE BLOOD IN KALA-AZAR

In the Indian Medical Gazette, 1917, and in the Treatise on Kala-azar by Brahmachari (2nd Edition, 1920) it was pointed out that the copious precipitate that was found when distilled water was added to kala-azar serum was globulin-like in nature and it was suggested that this might be due to an easily precipitable globulin or excess of globulin in kala-azar blood. We have since estimated the globulin content of the blood in some cases of kala-azar and the following results have been obtained. The diagnosis in each case was made by the presence of L. D. bodies in the splenic blood.—

First Case	
Globulın	. 19 per cent
Albumin	12 ,,
Second Case.	
Globulın	. 186 per cent
Albumin	1 03 ,,
Third Case	
Globulin	2 1 per cent
Albumin	172,,
Fourth Case	
Globulin	19 per cent
Albumin	1 36 ,,

In a series of healthy Indians the following results were obtained —

Total protein	49 to 69 per cent
Albumin	3 5 to 4 2
Globulin	1 0 to 2 7

The following figures have been obtained by other observers —

Howel

Total protein	6 014 to 7 6 per cent
Albumin	4 52 р с
Globulin	3 lpc

McLeod

Total protein	67 to 87 p c
Albumin	4 95 to 7 7 p c
Globulin	1 to 2 54 p c

Cholesterol content of the Precipitate obtained by Diluting Blood with Excess of Distilled Water

After diluting a volume of blood with 50 parts of distilled water and extracting the precipitate formed with ether alcohol chloroform and amyl alcohol the following figures were obtained —

rere obtained —	
Normal blood Kala azar blood	14 to 22 per cent
1st case	38 per cent
2nd	29
3rd	43
4th	46

Average 39 per cent

It will be seen that while, in kala-azar, the total protein and albumin contents of the blood are diminished, the cholesterol content in the precipitate obtained by diluting the blood with 50 parts of distilled water and the proportion of globulin to albumin contents are increased

We have observed that when kala-azar blood is mixed with distilled water, the precipitate formed shows the presence of many red corpuscles under the microscope. The precipitate, in our opinion, consists partly of globulin and partly of walls of red corpuscles. Though most of the red corpuscles present in the precipitate are nothing but shadow corpuscles, we have still to determine whether the excess of cholesterol present in kala-azar blood interferes with the hæmolysing effect of distilled water upon the red corpuscles in kala-azar blood and this will form the subject of a future investigation.

THE TREATMENT OF KALA AZAR WITH SOME NEW ANTIMONIAL PREPARATIONS

The new antimonial compounds which I am going to describe in this paper may be divided into two classes —

- (1) New antimonyl tartrates
- (2) Arvl or phenyl antimonial compounds
- (I) includes (a) urea antimonyl tartrate and (b) ammo nium antimonyl tartrate
- (2) includes (a) phenyl stibinate of sodium (b) acetyl p amino phenyl stibinate of sodium and (c) p amino phenyl stibinate of sodium or antimony analogue of soamin

I shall now give a report of my experience with some of these antimonial compounds in the treatment of kala azar. As most of these compounds have only been recently used by me in this disease the report must be regarded as a preliminary one

I (a) Urea Antimonyl Tartrate

This is a new compound. The method of its prepara tion was described by me at the July meeting of the Asiatic Society of Bengal 1920 and published in the Journal and Proceedings of the Society Vol XVI 1920. The amount of antimony present in it is nearly 38 per cent. It has been used by me both intravenously and intramuscularly. Intravenously it has up to now, been used in four cases.

The following are the notes of these cases -

(1) Patient K., æt 35 Leishman-Donovan bodies present in the spleen 2 to 5 c. c. of 2 per cent solution injected

Result of Treatment—Blood (1) Red blood corpuscles—2,400,000; white blood corpuscles—1,000, hæmoglobin—38 per cent on July 16, 1920, before treatment (2) Red blood corpuscles—3,000,000, white blood corpuscles—5,800, hæmoglobin—44 per cent, on November 13, 1920, after treatment Spleen extended 5½ in below the costal arch on July 16, 1920, before treatment, and 2½ in below the costal arch on November 13, 1920, after treatment Increase of body weight, 7 lbs Fever stopped after twelve injections No Leishman-Donovan bodies found in the spleen after fifteen injections

(2) Patient S, æt. 15 Leishman-Donovan bodies present in the spleen 2 to 4 c c. of 2 per cent solution injected

Result of Treatment.—Blood (1) Red blood corpuscles—3,600,000, white blood corpuscles—4,000; hæmoglobin, 46 per cent before treatment (2) Red blood corpuscles—3,900,000, white blood corpuscles—10,400, hæmoglobin—52 per cent, on November 1, 1920, after seventeen injections Spleen extended 5 in below the costal arch at the beginning of treatment and 1½ in below the costal arch after seventeen injections. No Leishman-Donovan bodies found on spleen puncture after completion of treatment Increase in body weight, I stone Fever stopped after eight injections

(3) Patient H, æt 20 Leishman-Donovan bodies present in the spleen 2 to 4 c c of 2 per cent solution injected

Result of Treatment —Blood (1) Red blood corpuscles—2,400,000, white blood corpuscles, 1,600, hæmoglobin—32 per cent before treatment (2) Red blood corpuscles—

3 500 000 white blood corpuscles—10 000 hæmoglobin—52 per cent on November 13 1920 after sixteen injections Spleen extended 5½ in below the costal arch before treatment and 1½ in below the costal arch after sixteen injections No Leishman Donovan bodies found on spleen puncture after completion of treatment Increase of body weight 2 lbs Fever stopped after three injections

(4) Patient K et 26 Leishman Donovan bodies present in the spleen 2 to 6 c c of a 2 per cent solution

injected

Result of Treatment —Blood (1) Red blood corpuscles—2 400 000 white blood corpuscles—1 200 hæmoglobin—38 per cent before treatment (2) Red blood corpuscles—2 700 000 white blood corpuscles—3 400 hæmoglobin—42 per cent after eleven injections Spleen extended 5 in below the costal arch before treatment and 4 in below the costal arch after eleven injections

Up to now urea antimonyl tartrate does not seem to be superior to tartar emetic or antimonyl sodium tartrate but it appears to me that symptoms such as vomiting severe cough or high rise of temperature do not follow the intra venous injections of urea antimonyl tartrate

Intramuscularly this preparation 'as up to now been used in two cases

The following are the notes of these cases -

(I) Patient S æt 24 Leishman Donovan bodies found on spleen puncture

Result of Treatment —Blood (I) Red blood corpuscles—I 400 000 white blood corpuscles—I 600 hæmoglobin—22 per cent before treatment (2) Red blood corpuscles—2,300 000 white blood corpuscles—4 800 hæmoglobin—32 per cent after eleven injections (3) Red blood corpuscles—3 300 000 white blood corpuscles—7 800 hæmoglobin—40 per cent after twenty injections Spleen 4! in below the costal arch before treatment and 3 in below

the costal arch after eleven injections and 1 in after twenty injections. Body weight same as before. No Leishman-Donovan bodies found on spleen puncture after twenty injections. Dose—1 to 2 grs. daily. There was some inflammation but never any suppuration at the sites of injection.

(2) Patient D., æt 10 Leishman-Donovan bodies present in the spleen

Result of Treatment —Blood (1) Red blood corpuscles—2,800,000, white blood corpuscles—3,800, hæmoglobin—30 per cent before treatment (2) Red blood corpuscles—3,600,000, white blood corpuscles—3,400, hæmoglobin—42 per cent after twelve injections. (3) Red blood corpuscles—3,600,000, white blood corpuscles—4,400, hæmoglobin—48 per cent after twenty injections Size of spleen reduced by 1 in Dose—½ to 1 gr Local reaction—no abscess but sometimes swelling and inflammation Fever stopped after sixteen injections. No Leishman-Donovan bodies found on spleen puncture after sixteen injections

In both the above cases the local reactions were less severe than what are met with in the case of potassium antimonyl tartrate.

(b) Ammonium Antimonyl Tartrate

This preparation has up to now been used by me only intramuscularly. The amount of antimony present in it is nearly 40 per cent. I have prepared this salt by neutralizing hyper-acid antimonyl tartrate with ammonium carbonate, and washing the precipitate with absolute alcohol.

The following are the notes of one case treated with it intramuscularly:—

Patient B., æt 10 Leishman-Donovan bodies present in the spleen. (1) Red blood corpuscles—1,600,000, white blood corpuscles—800, hæmoglobin—36 per cent before

treatment (2) Red blood corpuscles—2 400 000 white blood corpuscles—2 800 hæmoglobin—40 per cent after three injections Spleen 5 in below the costal arch before treatment and 3 in after three injections Dose—1 to 2 c c of 2 per cent solution Fever stopped after three injections Treatment still being continued

Intramuscularly this salt is less irritating than Tzuki s antiluetin which is aminonium potassium antimonyl tartrate

NB—It is interesting to note that the leucocyte count was so low as 800 before commencement of treatment

II Aryl Antimonial Compounds

I have been successful in preparing these compounds with the help of my chemist who has been working under me under a grant from the Indian Research Fund Association

The following compounds have already been made -

- (a) Phenyl stibinic acid and its sodium and ammonium salts
 - (b) p Amino phenyl stibinic acid and its sodium salt
- (c) Acetyl amino phenyl stibinic \mbox{acid} and \mbox{its} sodium salt
- (a) The salts of phenyl stibinic acid are too irritating and too toxic to be used for therapeutic purposes
- (b) The amino aryl compounds are extremely difficult to prepare Stibenyl which is allied to the acetyl compound has been used by me in two cases in one intra muscularly and in the other intravenously

The following are the notes of these cases The treat ment is still being continued

(1) Patient M , at 35 Leishman Donovan bodies found in the spleen (1) Red blood corpuscles—2 100 000

white blood corpuscles—2,800, hæmoglobin—32 per cent before treatment (2) Red blood corpuscles—2,600,000, white blood corpuscles—2,200, hæmoglobin—34 per cent after six injections. (3) Red blood corpuscles—3,400,000, white blood corpuscles—8,200, hæmoglobin—48 per cent after eleven injections linjections given intravenously on alternate days. After the ninth injection patient developed eruptions on his body similar to chicken-pox. Spleen could just be felt below the costal arch. Increase of body weight, 2 stone. No Leishman-Donovan bodies on spleen puncture. Doses—(1):1 grm., (2):15 grm., (3):2 grm., (4):3 grm., (5):4 grm., (6):5 grm., (7):6 grm., (8):8 grm., (9):1 grm., (10):1.5 grms., (11):2 grms.

(2) Patient H, æt 19. Leishman-Donovan bodies found in the spleen (1) Red blood corpuscles—2,900,000, white blood corpuscles—1,800, hæmoglobin—54 per cent before treatment (2) Red blood corpuscles—3,900,000, white blood corpuscles—2,800, hæmoglobin—50 per cent after five injections Doses—(1) 1 grm, (2) 15 grm, (3) 2 grm, (4) 3 grm, (5) 8 grm All the doses were given intramuscularly

The injections were given on alternate days There was much local irritation with pain and effusion into the injected parts which slowly subsided

This patient also developed eruptions similar to the above after the last injection

(c) p-Amino-phenyl-stibinate of sodium (antimony analogue of soamin) — This has been used by me in one case intramuscularly in 1-grain doses given every day. No local reaction. I propose to give it the name of Stibamine Subsequently I have been using it in bigger doses of 2 to 3 grm, as its toxicity appears to be low.

So far, it is too early to give any definite opinion about the effect of this antimonial preparation

The best antimonial preparation to be used in the treat ment of kala azar has not vet been discovered. Tartar emetic and antimonyl sodium tartrate have their serious drawbacks with which unfortunately we are more or less familiar The discovery of the amino antimony analogues of arsenical compounds opens up a new vista in the treat ment of the disease Anyone who thinks that the last word about the best antimony preparation has already been told in tartar emetic or antimonyl sodium tartrate is wrong One must pass from one antimony preparation to another to discover the one that is best It is stated that Ehrlich used nearly 600 arsenical preparations in 900 days from 1910 1913 in his attempt to discover the best one for the treatment of spirillosis And still we hear of newer arsenical compounds such as methylated salvarsan compounds hex amino arseno benzene triamino phenyl arsenic acid silver salvarsan and others Something approaching this has just been begun in the case of antimony

To me it appears that a day will come when in studying the organic derivatives of antimony one will be reminded of a simile employed by Dr. Bertheim about the chemistry of organic assenical compounds. He compares it to a sleeping beauty slumbering until quite recently in an unfrequented corner of Beilstein but who now awakened appears as one of the fairy gifts which synthetic chemistry bestows from time to time upon mankind. Let us who have to deal with dreadful kala azer hope that such a fairy will be discovered in the case of antimony. The words of Basil Valentine who stated centuries ago that he who deals with antimony must have an ample mind are very true.

May I incidentally refer here to the leucocyte increasing property of narcotine when given intravenously in solution in tartaric acid. This has already been noted by me

The two following cases of kala-azar illustrate the leuco-cyte-increasing properties of narcotine —

- (1) Patient M, æt. 12 Leishman-Donovan bodies present in the spleen (1) Red blood corpuscles—3,200,000; white blood corpuscles—3,400, hæmoglobin—42 per cent before treatment (2) Red blood corpuscles—3,400,000, white blood corpuscles 7,600, hæmoglobin—48 per cent after twenty-two injections Condition of spleen, same as before Increase of body weight—5 lbs. in three weeks. The increase of the leucocytes does not seem to be temporary Dose—1 to 1 gr, given every day
- (2) Patient A, æt 30. Leishman-Donovan bodies found in the spleen. (1) Red blood corpuscles—2,400,000, white blood corpuscles—2,400, hæmoglobin—46 per cent before injection (2) Red blood corpuscles—3,000,000; white blood corpuscles—5,200, hæmoglobin—46 per cent after twenty-six injections

I have been able to prepare a new compound of narcotine antimonyl tartrate as a definite crystalline substance. It is sparingly soluble in water but easily soluble in tartaric acid

Remarks

- (1) Urea antimonyl tartrate, a new definite compound, has been prepared and used in kala-azar intravenously as well as intramuscularly
- (2) Ammonium antimonyl tartrate has been prepared in a pure state—It is less irritating—than antiluetin—It has been used intramuscularly in kala-azar
- (3) Narcotine antimonyl tartrate, a new compound, has been prepared. Its leucocyte-increasing property has been noted
- (4) Acetyl-p-amino-phenyl-stibinate of sodium, which is allied to the patented Stibenyl, has been prepared and used in kala-azar.

(5) p Amino phenyl stibinate of sodium which is the antimony analogue of soamin has been prepared and is being used in kala azar—Its toxicity appears to be low—I propose to give it the name Stibamine

(6) As the best antimonial preparation for the treatment of kala azar has yet to be discovered, one must not rest con tented with the use of tartar emetic or sodium antimonyl tartrate

THE GLOBULIN OPACITY TEST IN KALA-AZAR

Two simple serum tests for kala-azar were described by Brahmachari (Indian Medical Gazette, December, 1917) and were named as (1) the globulin precipitation test and (2) the globulin ring test. The former consists in mixing one part of serum with two parts of distilled water when a distinct precipitate forms in the case of kala-azar serum. The latter consists in adding a few drops of distilled water on to the top of serum from a kala-azar patient when a distinct turbidity forms at the top of the serum. These tests have recently been confirmed by Milo, working in the University of Messina, and by some observers in China

That the above precipitate is a globulin is proved by the following tests —

- (1) It is soluble in normal saline, in dilute acids, in sodium bicarbonate solution and in sodium hydroxide solution
- (2) It is precipitated from its solution in normal saline when the solution is treated with equal parts of saturated solution of (NH₄)₂SO₄ or when it is saturated with MgSO₄ or NaCl.
- (3) It is not precipitated by NH4OH from its solution in dilute acids
 - (4) It is insoluble in distilled water

It has been subsequently found that if the globulin obtained by treating one part of serum with two parts of distilled water be dissolved in the serum of an individual on which formaldehyde has no action an opacity is obtained if a drop of formaldehyde is added to it If the same globulin is dissolved in normal saline it also gives rise to an opacity or precipitate when formaldehyde is added to the solution especially when the solution is rendered faintly alkaline by the addition of a little sodium bicarbonate This opacity however is generally less than that which is obtained when formaldehyde is added to the original serum lt is probably due to electrolytes other than sodium chloride being present in the serum. There is no doubt that this easily precipitable globulin which was described some time ago in kala azar serum by Brahmachari is responsibile for the aldehyde test

The Globulin Opacity Test

By estimating quantitatively the amount of water precipitable globulins present in a serum—we have succeeded in discovering a test which we propose to call the globulin opacity test for kala azar—The test is carried out as follows—

One part of serum is mixed with 6 parts of distilled water when a turbidity forms. The precipitated globulin after being uniformly mixed with the diluted serum is poured into a graduated cylinder the diameter of which is one inch. On looking through the height of the fluid containing the precipitated globulin over some black spots fixed to the bottom of the cylinder and adding more and more of the fluid till the spots become just invisible a point is reached which gives an estimate of the globulin precipitated

We have observed that in kala-azar a value is obtained which is fairly diagnostic of the disease, as will be seen in the following table.

Precipitated globulin (1 part of serum + 6 parts of H₂O) Height in inches at which the black spots disappear

Kala-azar				1 1
Do				14
Do				16
Do		•		0 7
Do				13
D_{o}				0 7 5
Do				1 25
Do				0.75
Do				09
Do				0 85
Do				09
Do				1 25
Do				1 25
Do				1.25
Malarial fo	ever			3 25
Do				3
Typhoid f	ever			35
Do				3
Pneumon	ıa			2
Nephritis				4 2
Aneurism				5 4
Hemipleg				5
Healthy s			above	8
D_{o}	Do	•	do	10
Do	Do		do	10
D_{o}	Do		do	10
Do	Do		do	10
Do	Do		do	10
Do	Do			6 5

NB—The height in inches at which the figures disappeared is *inversely proportional* to the amount of the globulin present in the serum

From the above we may conclude that if in any case the height of globulin precipitated by diluting the serum with 6 parts of distilled water and estimated in the above way is 1 25 in or less it may be regarded as fairly diagnostic of kala azar

We have also discovered that the total amount of water precipitable globulins present in kala azar is greater than that generally found in health or in other diseases. The total water precipitable globulins are obtained by diluting one part of the serum with 200 parts of distilled water and estimating it in the same way as above

The following table gives the value of the total water precipitable globulins in certain diseases

Preptdglobln(Iptofum+200 pits of H₂O) Hightin hit whih this kpt dippe

of	th blkptd ppe
Phthisis	3 1
Do	
	3 9
Do	3 15
Kala azar	19
Do	1 45
Do	16
Do	1 75
Do	19
Do	15
Do	18
Do	1 35
Do	1 65
D _o	14
D _o	1 65
Do	1 25
D _o	17
Do	1 35
Enlarged spleen (not leishmaniasis)	2 5
Dysentery	2 7
Do .	2 5
Chronic dysentery	5 5
Liver abscess with broncho pneumor	na 29

Precipitated globulin (1 part of serum+200 parts of H₂O) Height in inches at which the black spots disappear

Broncho-pneumonia with enlarged spleen	4 4
Broncho-pneumonia	39
Do	24
Do	37
Chronic bronchitis	35
Chronic rheumatism	4 4
Rheumatism	35
Mitral regurgitation	39
Influenza	37
Do	70
Do	7 0
Bright's disease	6 5
Pericarditis	63
Cancer	3 4

NB—The amount of the water-precipitable globulin is inversely proportional to the height in inches on the right hand of the table.

Conclusions

- (1) The easily precipitable globulins discovered some years ago are responsible for the aldehyde test. They are the same globulins that give rise to the globulin precipitation test and globulin ring test of Brahmachari
- (2) The total content of water-precipitable globulins is generally greater in kala-azar than either in health or in other diseases
- (3) A test has been described here, the globulin opacity test, for kala-azar By this quantitative test a more definite serum test is obtained than any hitherto known

UREA STIBAMINE IN KALA-AZAR

Proceedings of a meeting held at the Calcutta Medical Club on the 20th September 1923 Dr Upendranath Brahmachan read a paper on Recent Advances in the Antimomal Treatment of Kala Azar by the use of Urea Stibamine Sir Niltatan Sircar presided

Aastracts

The speaker said that the first series of his cases treated with urea stibamine appeared in the Indian Journal of Medical Re search in October 1922 Major Shortt's paper in the Indian Medical Gazette of July 1923 confirmed his observations as to the leishmanicidal properties of the compound Subsequent observations of Major Shortt were still more remarkable as three of his recent cases were eured respectively with 9 grm in 5 injections 75 grm in 5 injections and 65 grm in 4 injections In each of these cases the eure was established by subsequent observations in hospital and negative results obtained by culture of spleen puncture material In the present paper Dr Brahmachari was only limiting his observations on the effect of urea tibamine in the resistant or refractory cases By refractory or resistant cases of kala azar, he meant cases which had resisted treatment with two grams or more of sodium or potassium antimonyl tartrate given intravenously in the routine form of treatment extending over a period of 2 to 23 months or more It was known to every practitioner that a certain percentage of cases was not cured or sometimes not even benefited by antimonyl tartrates unless pushed for a very long time in some cases symptoms of intolerance to wards the drug would appear after administration of 2 grms or even less Some time ago certain observations conducted in Shillong led to the conclusion that all cases of kala azar were curable with 2 grms of tartar emetic. Subsequent observations did not confirm this view as a large majority of cases in Shortt's as well as Mackie's series were not cured with 2 grms of sodium antimonyl tartrate In Dr Brahma chari s series many cases were not cured by 2 grms but

about 10 per cent of the cases required 5-6 grms and about 5 per cent were absolutely refractory. Some of his cases had 6 grms and were not cured. He desired every practitioner to keep a record of the failures of the antimonyl tartrates. This would lead to the finding that the last word in the treatment of kala-azar had not been said in tartar emetic or sodium antimonyl tartrate. He observed that this fact as also the very prolonged course of treatment required in most cases justified the demand for further advances in the antimony treatment of kala-azar.

Dr. Brahmachari gave records of about a dozen of refractory cases with full details, first describing the effect of treatment with antimonyl tartrates and then reporting the results obtained after intravenous injections of urea stibamine Short notes of three of these cases are given below —

(1) Patient R-fever 6 months

Condition on admission—Temp 99—100° F. Spleen—extended 6 inches below the costal margin

Blood—R B C —3,000,000, W B C —3,500, Hb —40%

Peripheral blood culture and spleen puncture—positive

Patient was originally treated with antimonyl tartrates, 6 grms of sodium and 2 2 grms of potassium in 75 injections over 6 months. Symptoms of intolerance at times. Had soamin 2 grms in 11 injections and 6 T C C O, but to no effect. Patient went to Darjeeling and after 2½ months came back almost in the same condition—a refractory case.

Treatment with usea stibamine—2 grms in 9 injections starting from 15, increasing by 05, given twice weekly

After the third injection fever stopped

Condition after the injections—Fever nil, spleen just pal-

pable below the costal margin

Blood—R B C —5,000,000, W B C.—6,250, Hb — 70% Peripheral blood and splenic blood culture and spleen puncture—negative

Two and a half months after—Patient in excellent health, no fever, no enlargement of spleen, blood—normal

Patient cured

(2) Patient Mrs L-

Condition on admission—Fever 99—100° F. Spleen—extended seven inches below the costal margin Peripheral blood culture and spleen puncture—positive. RBC—3,000,000, WBC—2,400, Hb.—50%.

Patient was originally treated with potassium antimonyl tartrate -2 8 grms in 40 injections over a period of 6 months General condition worse there was loss of weight with fever of a low intermittent type Spleen extended 6 inches Cultural result and spleen puncture—positive A refractory case

Treatment with urea stibamine commenced one month after the last injection 16 grms in 9 injections in doses

from 1 to 25 grm during 6 weeks

Effect of treatment-One month after completion of treat ment-general condition improved Weight increased by half a stone No enlargement of spleen Fever stopped after 5 injections of the salt Cultural result and spleen puncture—negative
Blood condition two months after—Hb —60% RBC—

5 000 000 W B C — 7 800 Body weight increased by one stone Patient cured

(3) Patient N—

Condition on admission—Temp 100—103° F Spleen extended 5½' below the costal margin Blood Hb —26% R B C —2 500 000 W B C —1 000 Peripheral blood culture and spleen puncture—positive Cancrum oris and

cedema of the extremities

Patient was originally treated with sodium intimonyl tartrate—3 8 grms in 5 injections during 6 months. No improvement of the general condition no diminution of the size of the spleen loss of weight by 10 lbs cedema well marked cancrum oris diminished Temp 90 100° F Symptoms of intolerance after 2 8 grms

Blood-Hb -32% R B C -3 250 000 W B C -2 500 Culture and spleen puncture—positive A refractory case

Treatment with urea stibamine—2 85 grms in 13 injections extending over two months in doses of 1 to 25 grm

Effect of treatment—General condition—great improvement Body weight increased by one stone No enlargement of spleen Blood—Hb—54% RBC—4 250 000 WBC—6 200 Culture and puncture—negative Patient cured

The speaker observed that in most of his cases urea stibamine treatment was started about 11 to 2 months after the stoppage of the previous antimony treatment and as he reported in his paper on

"Chemotherapy of Antimonial Compounds in Kala-azai Infection" in the Indian Journal of Medical Research, that the excretion of antimony was complete by I to 1½ months, the question of the effect of residual antimony from the antimonial tartrates in the system did not arise in such cases. As to the purity of the antimonyl tartrates he stated that his salts were chemically pure and were specially made for purposes of research

All the cases reported were treated with intravenous injections. The speaker stated that both he and Major Shortt were also using the drug intramuscularly. The local reaction was little. But it was not yet time to dwell on the few cases treated intramuscularly though he expected very

encouraging results

He stated that the salt was being prepared in his research laboratory under trained experts and strictest supervision as well as with very strict aseptic precautions. It was repeatedly tested both for toxicity and sterility. The solution was not boiled before use

Dr Brahmachari then referred to another compound, stibamine a name given by him to the sodium salt of p-amino-phenyl-stibinic acid. This salt had the same relation to stibacetin as soamin or atoxyl has to ars-acetin. Its chemical and physical properties as also the toxicity had already been published in the Journal of Tropical Medicine and Hygiene, August, 1921 and in the Indian Journal of Medical Research, October, 1922. He reported three cases treated with stibamine, in two of which the result was satisfactory. The patients were cured with 2.4 and 1.2 grms respectively. In the third case fever was absent and blood condition improved. Culture and spleen puncture were both negative but the spleen still remained palpable.

Dr Brahmachari remarked that the duration of treatment and the number of injections required for complete cure with urea stibamine were less than with any other antimony preparations. His results had been confirmed by

Major Shortt's observations

In conclusion the speaker observed that-

(1) Urea stibamine was superior to all other antimony salts for the remarkably short course of treatment. Fewer injections were required for a complete cure

(2) Refractory cases yielded to urea stibamine very

strikingly.

(3) No cases had yet been resistant to this salt

(4) No relapse had yet come to his knowledge although some of the cured cases were under his observation for nearly two years after complete cure

(5) There is a possibility of its being used inframus

cularly as in the case of atoxyl or sormin

Dr Napicr after some preliminary remarks gave his experience about urea stibimine which was still very limited A boy 12 years old had been suffering from kala azar for four months His spleen was four inches and liver two inches below the costal margin WBC count was 3,000 He had 10 injections of urea stibamine and at the end of 4 weeks his spleen could not be felt. The important point about the case was that after two weeks his W B C count was over 6 000 Dr Napicr considered the case com pletely cured One or two points he would however like to speak on the subject of treatment on Dr Brahmachari s paper Dr Brahmachari quoted Major Shortt s experience Shillong Personally he (Dr Napier) was absolutely convinced about the effect of urea stibamine It was possible that the strikingly good results obtained by Major Shortt with urca stibimine were to some extent due to the climatic con dition in Shillong His experience about the treatment of kala azar with sodium antimonyl tartrate was that 2 grms of the salt were not always sufficient in bringing about a complete He observed that in Calcutta very small number of cases were cured by two grams Dr Brahmachari s cases could be considered as cured because after the lapse of a year he found that they were in good health Dr Napier did not consider that negative results of spleen puncture and blood culture were the last words on the subject of cure He knew of two cases which gave positive cultural results after a course of treatment with sodium antimonyl tartrate but were subsequently found to have been completely cured after one year without any further treatment

Dr Umaprasanna Basu after referring to his experience in the treatment of kala azar with sodium intimonyl tartrate in his general practice and in consultation with Sir. Leonard Rogers in two of his cases was of the opinion that sodium antimonyl tartrate was not the specific in kala azar in the proper sense of the term. In his experience out of 30 or 4 cases only 6 or 7 cases were cured. It took a very long time and then symptoms of intolerance were remarkable.

it could not be a specific and the researches of Dr Brahmachari had heralded its advent by the discovery of a true specific in urea stibamine. He thought that all were certainly grateful to Dr Brahmachari for the research he had carried out. The results were very encouraging indeed

C Sengupta said that in cases of kala-azar the impression that one forms of sodium antimonyl tartrate was that a large number of cases did very well but every case took a very long time. On the other hand in many cases the spleen did not diminish size, and R B.C as well as W.B C as also Hb did not seem to increase. Then some of these cases showed intolerance to the drug and some developed broncho-pneumonia, dysentery or diarrhœa during treatment with sodium antimonyl tartrate. He had observed that a certain number of cases which did not do well with sodium antimonyl tartrate, subsequently seemed to do well when potassium antimonyl tartrate was used Dr Brahmachari stated elsewhere, that cases which did not do well with sodium or potassium antimonyl tartrate treatment did well with metallic antimony. His own impression was that antimony was really the specific for kala-azar But as to the salt which should be used, of course we could not do better than to leave the question to Dr Biahmachari to solve He did not know the reason why a particular case did very well up to a certain limit and why it was that it did not do well any further. His criticism was only an enquiry whether they should be satisfied with urea stibamine as a salt pre-emmently specific for kala-azar He asked whether the variable solubility of a particular antimony salt got anything to do with the destruction of the parasite

Dr J M Das said that he had seen that sodium antimonyl tartrate in some cases was quite useless. Sometimes actual danger was apprehended. He observed that if in any way the WBC could be increased then antimonyl tartrates did well. He was trying in his own way to increase WBC and he got very good results. He gave antimony and tried to increase the leucocytes, by bringing on an artificial inflammation by injecting several substances along with sodium antimonyl tartrate. He suggested different things to the mofussil practitioners including the stem of nim as "gool" under the skin and into the muscle. While sodium antimonyl tartrate absolutely failed,

this process generally brought on good results. In his experience where sodium antimonyl tartrate failed potassium antimonyl tartrate did well. Hence before they passed any remarks against the antimonyl tartrates they should consider them more leniently and from a wider point of view.

Dr Satya Saran Mitra said that from the observations made by Dr Brahmachari he noticed that the swing of the pendulum had been turned against tartar emetic and sodium antimonyl tartrate and some other antimonial compound was making a rapid headway pushing them to the background People were getting accustomed to see only one side of the thing and they thought that sodium antimonyl tartrate was really the panacea although it was sometimes producing very disastrous results Now that some kind of check was put on the swing of the pendulum people would be on their guard in the u e antimonyl tartrates He asked Dr Brahmachari to express his opinion and observations on the use of the new drug urea stibamine by the intramuscular route and whether he found it to be efficacious in the treatment of many diseases which were as fell as kala azar such as the infantile cirrhosis of the liver He wanted to know the circumstances that led Dr Brahmacharı to the discovery of the new drug

Dr Bepin Behari Gupta said that this lecture was heard by him in the Asiatic Society of Bengal. It was glorious that a Bengalee made this most important advancement in the treatment of a disease which was really black black in its treatment black in all its aspects and that blackness was going to be whitened by one of Bengal's own men

DR BRAHMACHARI S REPLY

He discussed the various points raised by the gentlemen present and said

First of all I must thank Dr Napier most heartily for his remarks I do not think that the good results obtained by Major Shortt in Shillong were to any great extent attributable to good climate. If we go through his last paper that was published in the Indian Medical Gazette we will find that there were resistant cases cases which were resistant to the antimonyl tartrates even in Shillong inspite of the best climate possible there. Of course climate to some extent may have some influence but there was something more in urea suba mine that was responsible for the cure of cases of kala azar in such a remarkable way as recorded by Major Shortt.

"As regards complete cure, I had the patience of observing a series of cases kept in the Campbell Hospital for about 2 years for observation The patients are still there and are perfectly cured. There is no enlargement of the spleen, blood count is normal and the culture from spleen puncture as well as from peripheral blood is negative. There does not exist any sign of the disease in these patients course the definition of cure of a disease must be very difficult If, however, one or two years have passed and no symptoms and no trace of the disease have been found. the blood count is quite normal, there is no anæmia and the blood culture and spleen puncture are negative, I do not think one should question their cure Although we cannot possibly claim that the last word about the treatment of kala-azar has been said in urea stibamine, I hope that the discovery of this salt has advanced greatly the treatment of the disease At least we can say that the antimonyl tartrates were not the last words. There is no doubt that research must be carried on along proper lines so that we can bring out, for the treatment of kala-azar specifics like salvarsan or neo-salvarsan. Urea stibamine being allied to soamin or atoxyl is a step towards the discovery of such compounds From the study of atoxyl I felt that the corresponding urea salt of antimony might be very suitable for the treatment It also struck me that being an urea ester, it of kala-azar would be more suitable for intramuscular injection as urea is a local anæsthetic This was the origin of urea stibamine and my speculations have been justified by actual experience There is no doubt that the cases read before the meeting this evening distinctly prove that we have made a very great advance in the treatment of kala-azar by the use of urea stibamine.

"It cannot be said that insufficient doses of the sodium or potassium antimonyl tartrate were always responsible for making cases refractory My cases were really very resistant

and I think you will agree with me in this point

"As regards the value of intramuscular injection of urea stibamine I cannot say anything definitely till after some more experience with the drug I am, however, glad to say that I met with success in two cases in which urea stibamine was used intramuscularly without any local reaction or pain. But more cases must be studied before we come to a definite conclusion about the intramuscular use of urea stibamine.

"As regards the advantage of alternating sodium and

potassium antimonyl tartrates. I hold that it really exists in some cases, but in the refractory cases described in my paper

this evening this was not so

'As regard Dr Mitra's remarks that the swing of the pendulum has gone back the other way I agree with him that the swing has gone back so far as the antimonyl tartrates are concerned but otherwise it has got a fresh momentum and the swing is always going forward Many of us once thought that sodium antimonyl tartrate was the specific in every case of kala azar but it is not so

As regards the supply of urea stibamine it is certainly limited 1 prepare urea stibamine in small quantities and 1 am giving it a trial in a scientific way 1 cannot say

whether I can at present supply larger quantities

It now remains for me to express my thanks to the Indian Research Fund Association for their valuable assist ance in carrying on my researches

PRESIDENT S REMARKS

I think I voice the sense of the whole house when I say that we offer our heartiest congratulations to Dr Brahmachari on this occasion. Ours are double congratulations. One is for the discovery of this urea antimony salt and the other for the application of that salt to the treatment of kala agar.

It is scepticism that is the basis of all scientific progress in this world It simply means that whenever any evidence is put before us it should be properly examined and then ac cepted for unless it is so done it is likely to crumble in the long run Well applying this principle in this particular case it is our duty to gather all information in regard to this matter with which we are so deeply concerned I mean the treatment of kala azar Dr Brahmachari adduces evidence of the strongest character that o far urea stibamine is the best agent for the treatment of kala azar in his and Major Shortt's hands Well we accept it so far as is concerned to himself or Major Shortt But to say that it has very greatly advanced the treatment of kala azar I would take a little time and I think Dr Brahmachari would not grudge me the time I want, because if after a couple of years we say that it is a great advancement in the treat ment of kala azar I think that word will be really and truly

said My suggestion is this, that a very good prima facie case has been made out on the evidences put forward by Dr Brahmachari's experience as well as that of Major Shortt in Assam So far as they go they are indisputable. Nevertheless we think these findings may have limitations A few years ago, it must be admitted, great expectations were entertained about the treatment of this fell disease with sodium antimonyl tartrate I do not think that hope has crumbled into dust to-day. But every case cannot be expected to be successful In course of time some of our apprehensions have been realised We hope we should not have a similar experience in regard to this new salt lt is possible that in the course of a few years our indefatigable worker will have again to draw our congratulations upon the discovery of yet a better and more efficient drug for the treatment of this disease I think that considered from the chemical, physiological and pharmacological points of view so far urea stibamine has stood the test in his own and Major Shortt's hands and I believe, it will stand the test with many of us

"Whatever the future may prove, even if we find out some improvement in the treatment of kala-azar, we shall always remain proud of Dr Brahmachari At this moment this salt comes as a real friend to enable to help many of our brothers and sisters to be cured of this fell disease, and we can surely realise how great is our delight to be able to depend upon one of our home-made products for the purpose of vanquishing one of our worst enemies No German would have stronger reasons of being proud of the Krupp gun nor a Frenchman would have greater reasons of being proud of his navy than we Bengalees would have, being able to use one of our home-made products for the purpose of fighting out one of the most dreadful diseases through the discovery of our Dr Brahmacharı We have every reason of being proud of what has come out of Dr Brahmachari and I believe that in the future we shall have occasion to hear from him fuller accounts about his work in the treatment of this fell disease?

THE RELATION BETWEEN THE CHEMI CAL CONSTITUTION OF ANTIMONIALS AND THEIR THERAPEUTIC PROPERTIES

In this paper the relationship of the chemical constitution of the following antimonials to their therapeutic properties are discussed —

- (1) Metallic antimony
- (2) Antimony trioxide
- (3) The antimonyl tartrates
- (4) Aromatic antimonials—urea stibamine

Antimony belongs to the odd series of Group V of Mendeleeff's periodic system of elements in which the gradual transition from typical non metals to typical metals is clearly exhibited Phosphorus is decidedly a non-metal while antimony and bismuth are typical metals although they are brittle Arsenic which stands between these two classes shows properties belonging to both groups of elements The acid producing properties of antimony are greater than those of bismuth and less than those of arsenic It has the property of combining with tartaric acid and giving rise to an acid of the type of what has been termed antimonyl tartaric acid Tartar emetic and sodium antimonyl tartrate should not be regarded as antimony salts of an organic acid They are really potassium and sodium salts of antimonyl tartaric acid as has been proved by Clarke Stallo Jungfleisch Guntz Adam and others Antimony exists in them not as a basic-Sb=O, in combination with

tartaric acid, but as ortho-antimonious acid, Sb $(OH)_3$, in which two of its hydroxyl groups are replaced by the divalent group $C_4H_4O_6$ giving rise to antimonyl tartaric acid. In other words, antimony in tartar emetic and allied compounds exists in an acidic state

The most important factor upon which the therapeutic value of an antimonial depends, is its property of containing trivalent antimony in an acidic state or its ability of being converted into a compound of this kind after its introduction into the body which will further possess the mobility of being converted into a compound containing the radicle -Sb=O in a reactive state or in a highly dispersed condition

The reactive state corresponds more or less to the nascent state of elements and mobility means the quickness with which the property referred to above is displayed

The antimonyl tartrates, finely divided metallic antimony or an aromatic antimonial derived from stibanilic acid, more or less possess these qualities and the superiority of an antimonial over another depends upon the degree of its power of exhibiting them.

In studying trypanosomiasis, Ehrlich held the view that trypanosomes assimilated the organic derivatives of arsenic only when the arsenic was present in the trivalent and not in the pentavalent form. Similarly the experiments of Kolle, Hartoch, Rothermundt and Schurmann have shown that compounds containing pentavalent antimony were not organotropic except in large doses and were, at the same time, slightly parasitotropic. Preparations containing trivalent antimony were, as a rule, toxic to the organism and at the same time it was also shown by these observers that for antimony compounds, soluble or insoluble, organic or inorganic, to be of therapeutic value in trypanosomiasis, the antimony must be in the trivalent form. My most recent researches and those of others that have followed

me have however proved that the aromatic pentavalent antimonials are much more potent in the treatment of kala azar than the trivalent antimonyl tartrates. I hold that it is not so much whether an antimony compound is trivalent or pentavalent that is responsible for its therapeutic value but its capacity for being quickly converted into a compound containing —Sb = O

By studying the excretion of antimony in man after intravenous injection of the aromatic antimonials of the type of urea stibamine and also of the antimonyl tartrates one can explain the superiority of the former over the latter on the above theory. I have observed that in the case of tartrar emetic the curve of excretion is one slowly converging to the base line.

The amount of antimony excreted in the urine during the first 24 hours after intravenous injections of tartar emetic is about 6 per cent of the amount injected The amount of antimony excreted in the urine during the first 24 hours after intravenous injections of urea stibamine is 30 to 40 per cent of the amount injected The excretion of antimony after intravenous injections of a pentavalent organic antimonial follows a curve the first portion of which representing the excretion during the first 24 hours is abrupt and the second portion follows a course similar to that found in the case of tartar emetic. It is pro bable that a pentavalent organic antimonial is converted in the body into a trivalent antimonial and that as long as it exists in the body in the pentavalent form its rate of excretion is much quicker than when it is converted into the trivalent form During the latter stage the curve of excretion is similar to that of tartar emetic in which antimony exists in the trivalent form. Since a great portion of antimony present in an aromatic pentavalent antimonial (urea stibamine) is quickly eliminated the chances of toxic action of the compound are much less than that of an antimonyl

tartrate. In the process of conversion of an aromatic pentavalent antimonial in the body into a compound containing trivalent antimony, a reactive—Sb=O, is formed, which is probably responsible for the remarkably beneficial results observed in the treatment of leishmaniasis by the use of unea stibamine

Finely Divided Metallic Antimony

Though various metals have been administered intravenously in the colloidal state, metallic antimony is perhaps the only one which has been put into the circulation in the crude form of a fine suspension. To Plimmer and Fry belongs the credit of first demonstrating the possibility of introducing metallic antimony into the veins without danger of capillary blocking. To Ranken belongs the credit of using the drug successfully in man in the treatment of trypanosomiasis by the intravenous route.

In 1915 I described a number of cases of kala-azar treated successfully with intravenous injections of metallic antimony and I pointed out that it was the most powerful leishmanicide that was known at that time, just as it was the most powerful of the known antimonial trypanocides

I observed that in cases in which the soluble salts of the type of tartar emetic did not show any improvement in the blood condition or temperature of the patient after several injections, finely divided metallic antimony administered intravenously brought about complete cure. In addition, the number of injections required for a course of treatment with metallic antimony was much smaller than those required in the case of the antimonyl tartrates. Three or four injections frequently cured the patient, though sometimes the injections required were as many as eight or nine. Even then the number of injections required for cure was less than what was generally required in the

case of antimonyl tartrates The only objection is the complicated technique of the operation of injection which is a serious obstacle to mass treatment of the disease

The mechanism by which metallic antimony is taken up into the system after intravenous injections is very interesting. Quickly taken up by the leucocytes and perhaps also by the cells of the reticulo endothelial system and without causing any capillary blocking it is converted into a soluble antimony compound as the particles of antimony sooner or later disappear from the leucocytes. I consider that it is converted into a compound in which the antimony exists in a trivalent state and this conclusion has been arrived at by me by following the curve of excretion of antimony in the urine after its administration which resembles the curve of excretion of antimony after administration of tartar emetic.

This trivalent antimony compound is subsequently converted into one containing a radicle of -Sb=O in the reactive stage or in a highly dispersed condition. It does not rest at the stage of a trivalent antimony compound allied to $Sb\ O$ because $Sb\ O_a$ when injected intravenously does not exhibit such therapeutic properties as those of metallic antimony as will be presently seen. It is possible that it is finally converted into nascent metallic antimony

Levaditi has propounded a general law with reference to all the members of the nitrogen family of elements occupying Group V of Mendeleeff's periodic table such as arsenic antimony or bismuth. They or their compounds exhibit their parasiticidal properties only after they have been acted upon by the tissues. If fresh extract of liver is added to them then they exert their parasiticidal properties.

I have observed that if a solution of tartar emetic or urea stibamine is mixed with a culture of the flagellated form of leishmania and the mixture examined under the microscope they do not die. Like bismuth or its compounds they become active only after they have been acted upon by the tissues

It has been suggested that in the case of bismuth, the action of the cellular extract gives rise to a new compound, 'bismoxyl,' and it is this which possesses the destructive power against the Tieponema pallidum. The substance in the extract which has the property of changing bismuth into bismoxyl, has been termed 'bismogene'

Bismoxyl is supposed to be a bismuth toxalbumin

Chemically, some of the bismuth compounds contain the radicle $-B_1=O$, just as some of the antimony compounds contain the radicle -Sb=O, and it is very likely that the bismuth toxalbumin also contains the radicle, $-B_1=O$, in the reactive stage, or in a highly dispersed condition. A corresponding antimony compound, which may be called 'stiboxyl, is probably formed in the case of antimony

It has been recently observed by Meleney that in kala-azar, clasmatocyte tissue is developed as a tissue reaction and probably, as I have suggested, out of the reticulo-endothelial system. I hold that this reticulo-endothelial system gives rise to the production of bismoxyl or stiboxyl as the case may be.

I have discussed in detail what, I consider, is the mechanism by which metallic antimony exerts its parasiticidal properties, because, being a simple element, it does not contain any groups or radicles which may complicate any explanation that may be suggested. The sequence of events in this mechanism may be summarized as follows—

Metallic antimony—taken up by leucocytes and cells of the reticulo-endothelial system—a soluble trivalent antimony compound—an antimony compound containing —Sb=O in the reactive stage or dispersed condition (stiboxyl), or nascent antimony which acts as a leishmanicide in kala-azar.

Colloidal Metallic Antimony

Colloidal metals are remarkable in having minimum organotropic properties and at the same time are frequently

parasitotropic. An ideal medicament should be one in which the ratio of dosis curativa to dosis tolerata should be as low as possible. Because of the extreme division of the metals in the colloidal state there is an immense surface of contact between a colloidal solution or suspension and the surrounding medium. For instance it has been calculated that the total surface of particles of gold in one cubic centimetre of colloidal gold may attain to nearly 6 500 square feet. This immense contact surface of the colloidal suspension of metals their electric charge of constant sign for the same substance and the fact that in the living body the reactions that take place are nearly always between colloids render the potentialities of metallic colloids very great.

Sb₂O₃-Antimony Trioxide

Yorke and Blackmore have used trixidine in oily suspen sion intramuscularly. They have also used a fine prepara tion of the same intravenously.

Kolle Hartoch Rothermundt and Schürmann consider that the formation of a deposit of an insoluble slowly absorb able compound of antimony such as antimony trioxide acts prophylactically against trypanosome infection. The principle of the employment of insoluble organic compounds of antimony either in ointment form or through the formation of intramuscular depots constitutes what the authors designate therapia mite curans as contrasted with therapia magna sterilans. Rogers used it in kala azar. In my experience it is weak in its therapeutic properties in kala azar.

I shall now try to explain why antimony trioxide is feebler in its leishmanicidal properties than metallic antimony. Though antimony exists in it in the trivalent state yet its proneness for being converted into a compound con taining $-\mathrm{Sb} = \mathrm{O}$ in a reactive state is slight as it is a fairly stable compound. This theory agrees with the fact that it is

more potent than Sb₂O₅ which is more stable and has much less leishmanicidal properties. Of all the oxides of antimony Sb₂O₄ is the most stable. So far as I am aware, Sb₂O₄ has no use whatever in therapeutics. One may, therefore, lay down as a general rule that the more stable an oxide of antimony is, the less is its trypanocidal or leishmanicidal property.

Tartar Emetic and other Antimonyl Tartrates

As stated before, these are salts of antimonyl tartaric acid and have been erroneously considered as antimony compounds of tartaric acid even in a recent textbook on kala-azar

Among the antimonyl tartrates of the type of tartar emetic or sodium antimonyl tartrate may be mentioned ammonium antimonyl tartrate, urea antimonyl tartrate, aniline antimonyl tartrate, ethyl antimonyl tartrate, quinine antimonyl tartrate, cinchonine tartrate and narcotine antimonyl tartrate

If T (NH_4) , T (Urea), T (K), T (Na), T (Aniline), etc , represent the toxicity of the above tartrates respectively, l have observed —

$$\frac{T \cdot (NH_4)}{T \cdot (Urea)} = \frac{T \cdot (NH_4)}{T \cdot (K)} = \frac{T \cdot (NH_4)}{T \cdot (Na)} = \frac{T \cdot (NH_4)}{T \cdot (Na)} = \frac{55}{60} \text{ or } \frac{11}{12}$$

If T Sb (NH₄), T, Sb (Urea), T Sb (K), T Sb (Anılıne), T Sb (Na) represent the toxicity of the antimony content of the above tartrates, we have —

$$\frac{T \text{ Sb(NH_4)}}{T \text{ Sb(Urea)}} = \frac{41}{46}, \quad \frac{T \text{ Sb(NH_4)}}{T \text{ Sb(K)}} = \frac{40}{46}, \quad \frac{T \text{ Sb(NH_4)}}{T \text{ Sb(Na)}} = \frac{38}{46}, \quad \frac{T \text{ Sb(NH_4)}}{T \text{ Sb(Aniline)}} = \frac{35}{46}$$

Therefore, in the case of the guinea-pigs, ammonium antimonyl tartrate is the least toxic, then comes the urea salt, then the sodium and potassium salts which are equally toxic and then the aniline salt

The maximum tolerating capacity of the same species of animals for a drug is directly proportional to its maximum tolerated dose

We thus have -

	Maximum with antim			of guinea	pigs treated = K ¹ × 03
/2\	D.	n n	f I		

(2)	Do	Do	Do	Urea antimonyl tartrate = K ¹ × 025
(3)	Do	Do	Do	Potassium antimonyl tartrate

= K¹ × 025

From this we conclude that of all the antimonyl tartrates used in the case of the guinea pigs their maximum tolerating capacity is with ammonium antimonyl tartrate and that the presence of N in the basic radicle of an antimonyl tartrate diminishes the toxicity of some of them

Generally speaking the toxicity of the antimonyl tartrates depends upon their antimony content. A notable exception is in the case of quinine antimonyl tartrate in which the toxicity is low. The possibility of using the compound in therapeutics should therefore be borne in mind as it may combine the therapeutic properties of antimony and quinine

I have not however been able to confirm the observations of Farghar and Gray that the toxicity of the antimony content of quinine antimonyl tartrate is only one fifth that of tartar emetic though I agree with them that its toxicity is less than that of tartar emetic I confirm their observations that quinine antimonyl tartrate on boiling with antimony trioxide is converted into the more toxic quino toxine antimonyl tartrate I have not been able to confirm their conclusions that the sodium salt is less toxic than the potas sum salt I have confirmed Plimmer and Thompson s

observations that the lithium salt is more toxic than the sodium or potassium salt and that the toxicity of the sodium and potassium salts is equal

I have further found that ammonium antimonyl tartrate is the least toxic of all the inorganic tartrates, the presence of nitrogen in the basic radicle diminishing its toxicity. Because of the high antimony content of the ammonium salt, its relatively low toxicity for the lower animals and likewise for human beings, and since it was found to possess a marked degree of therapeutic activity in the treatment of kala-azar, I consider it superior to both potassium and sodium antimonyl tartrates

I found that after the administration of a toxic dose of an antimonyl tartrate, the pathological changes are most marked in the lungs, kidneys, liver, pituitary and suprarenal glands, consisting chiefly of hæmorrhages into the substance of these organs and destruction of their cellular elements. Similar changes were produced by toxic doses of the new aromatic organic antimonials.

Delayed Antimony Poisoning —Cases of death in guineapigs three weeks or so after one injection of an antimonial salt have been met with, showing definite symptoms of antimony poisoning and presence of antimony in the viscera

These cases of delayed antimony poisoning are of very great clinical importance, as they prove that the excretion of the drug may sometimes be very slow after injection of antimonial compounds and some of the cases of sudden death during antimonial treatment may be due to a cumulative action of the drug

Cumulative and Tolerance Experiments with Tartar Emetic —I have observed that repeated injections of tartar emetic in sub-lethal doses did not give rise to any tolerance towards the drug except very rarely Generally the results pointed to a cumulative action of the drug, or at least made the animal susceptible to the next higher dose.

I have observed before that antimony in tartar emetic and other antimonyl tartrates exists in the form of antimoni ous acid Sb(OH), in which two hydroxyl groups have been replaced by the divalent C,H₁O₆ When introduced into the system its therapeutic value depends upon its ability to give rise to a reactive -Sb=O which, theoretically speaking should be the same as that of the salts of hydrated Sb 0, 1e Sb 0,+3H 0 or Sb(OH), or ortho antimonious acid Herein lies the superiority of the promatic antimonials over the antimonyl tartrates which we shall presently see On the other hand if it were possible to prepare an anti monial having the same composition as the antimonyl tartrates but having the radicle -Sb=O as is shown in the old configuration of tartar emetic and allied salts then such an isomer of tartar emetic would be more potent in the treatment of kala azar than tartar emetic itself. We await the production of such an isomer

Besides the antimonyl tartrates already referred to the following amino antimonyl tartrates have been prepared in my laboratory —

- (1) Phenocoll antimonyl tartrate
- (2) Anæsthesin antimonyl tartrate
- (3) Novocaine antimonyl tartrate
- (4) Aposthesine antimonyl tartrate
- (5) Orthoform antimonyl tartrate
- (6) Acriflavine antimonyl tartrate

The late Sir Patrick Manson once wrote to me as follows 'Go on in your efforts to get an antimony compound that can be used as an intramuscular injection or better still as a drug that can be administered by the mouth

The therapeutic value of an antimonial depends upon its concentration in the tissues after administration

Unfortunately ordinary antimonials cannot be administered orally, intramuscularly or per rectum in such doses as to bring about this concentration without at the same time giving rise to local distressing symptoms. The above new amino-antimonyl tartrates containing radicles, possessing anaesthetic properties, may be worth trial by these routes

Ointment of metallic antimony in a state of finest subdivision may be more easily absorbed and less irritating than that made with ordinary metallic antimony, and may be of therapeutic value in the treatment of kala-azar

Aromatic Antimonials

Let us now pass on to the consideration of the aromatic antimonials and their value in the chemotherapy of antimony.

In 1920, shortly after I had been financed by the Indian Research Fund Association for carrying on researches into the treatment of kala-azar, I brought to the notice of the Government and the Governing Body of the Indian Research Fund Association the possibility of the potentialities of organic antimonials in the treatment of Indian kala-azar, my conclusions being based on theoretical grounds, from an analogy of the value of the corresponding compounds of arsenic, namely, ars-acetin and atoxyl, in the treatment of certain protozoal diseases

The acetyl compound (stibacetin, stibenyl) was used more or less successfully outside India in the treatment of kala-azar and other forms of leishmaniasis (Caronia, Kharina-Marinuchi, Spagnolio) Manson-Bahr successfully used it in a case of kala-azar Early in 1921, I discovered that urea could combine with stibanilic acid and that the resulting compound surpassed all my expectations in its value in the treatment of kala-azar The discovery of this compound and my researches into the chemotherapy of antimonial com-

pounds in kala azar infection opened up a new vista in the treatment of the disease

The starting material of aromatic antimonials is acetyl p amino phenyl stibinic acid. Theoretically speaking the value of the sodium salt of the acid in the treatment of kala azar should be the same as that of ars acetin in the treatment of trypanosome infection. Are acetin has certain marked advantages compared with atoxyl being more stable and less toxic to some animals while equally toxic to the parasites. This diminiution of toxic effect is however noticeable only in certain animal species and not with horses or guinea pigs. Voegilin and Smith have observed that it is considerably less toxic than atoxyl and more trypanocidal, possessing a chemo therapeutic index about five times higher than atoxyl.

It is a well known theory in the case of aromatic arsenicals that their therapeutic value depends upon the reduction products produced after their introduction into the system. These reduction products probably all contain the reactive—As=O. The trivalent aromatic arsenicals of the arseno benzene group possess the property of producing these reduction products to a greater extent than the pentavalent arsenicals and hence their superiority in the treatment of treponema and trypanosome infections over the pentavalent arsenicals except tryparsamide. I hold that the therapeutic value of the aromatic antimonials also depends upon the same property.

The comparative value of the aromatic antimonials in the treatment of kala azar also depends upon their toxicity and parasitotropic properties following their administration. These again depend upon their chemical configuration and physico chemical properties. In order that they may be of their dosis curativa to that of their dosis tolerata must conform to Ehrlich's formula which is 1 3 or less.

Aromatic Antimonials of the Stibino-Benzene Group

Antimonials of the stibino-benzene type have not yet come into use in the treatment of human diseases, though they have been used with indefinite results in the case of certain diseases of animals.

Trivalent aromatic antimonials of the type of salvarsan or neo-salvarsan will probably be in future the highest advance in the antimony treatment of kala-azar.

Aromatic Antimonials derived from p-Stibanilic Acid (p-Amino-phenyl-stibinic acid)

Acetyl-para-aminophenyl-stibinate of sodium (Stibenyl, Stibacetin, Sodium acetyl-p-stibanilate)

The minimum lethal dose of phenyl stibinate of sodium, is three and half times less than that of acetyl-p-aminophenyl-stibinate of sodium, while its maximum tolerated dose is 35 times less. Injected into lower animals, it gives rise to hæmorrhagic nephritis and other symptoms of severe antimony poisoning. This compound has little or no use in therapeutics, but the introduction of NH₂ into its benzene nucleus at once diminishes its toxicity and raises its therapeutic value to a remarkable extent.

The acetyl compound of antimony has been used in the treatment of kala-azar but with unsatisfactory results. Besides, as has been shown by me, stibenyl becomes toxic with age in India and it has now come into disuse. But I still hold that pure acetyl-p-amino-phenyl-stibinate of sodium should again be given a trial in kala-azar and may in future be found to be free from all those toxic effects that were exhibited by stibenyl.

The sodium salt formed after hydrolysis of the acetyl compound corresponds to atoxyl or soamin and is sodium-p-stibanilate. Comparing its toxicity with that of the acetyl

compound, it will be seen that the introduction of the acetyl group into it does not reduce its toxicity as in the case of the corresponding arsenic compound. Thus while in the case of ars acetin the toxicity is markedly diminished by the introduction of the acetyl group into atoxyl being 1 that of atoxyl in the case of sodium stibanilate and the acetyl compound my observations have shown that their toxicity is the same. The M L D is 0.7 grm per kilo of body weight and the M T D is 0.35 grm per kilo of body weight in guinea pigs given intramuscularly in the case of both the compounds.

The pure salt is fairly stable. It has been stated by some observers in India that the compound is very easily decomposed. Evidently the substance that they were using was impure or not properly prepared. Three cases have been treated by me with this compound with satisfactory results. But as the number of cases was limited no attempt can at present be made to give a comparative estimate of the therapeutic values of sodium p stibanilate and urea stibamine—a compound to be discussed later on

Chloro Stibacetin (von Heyden 471) or Stibosan

This is a compound formed by the replacement of one hydrogen atom in the benzene nucleus of the acetyl compound by chlorine. The published results of cases treated with this compound lead to the conclusion that it is weaker in its therapeutic effects when compared with urea stibamine. It has been claimed that the introduction of chlorine increases its stability. It has also been claimed that it can be stored in ordinary stoppered bottles and weighed out when required and is therefore most useful for general purposes. In my opinion such a compound has more or less the same stability as the antimonates and therefore there is less chance of the production of the reactive.—Sb=O in the tissues after

their administration which, I consider, is responsible for the beneficial results following the administration of an antimony compound. This explains why antimonates in which antimony exists in a pentavalent form are of very little use in therapeutics, as they are very stable and quickly excreted unchanged after administration. The same also holds good in the case of arsenic.

Urea Stibamine

The next aromatic antimonial discovered by me is urea stibamine I shall discuss its therapeutic value later on.

Benzene-sulphon-p-amino-phenyl-stibinate of Sodium

The next aromatic antimonial of probable therapeutic value that has been discovered by me is benzene-sulphon-p-amino-phenyl-stibinate of sodium. The corresponding arsenic compound is known as hectine which possesses certain therapeutic properties in the treatment of syphilis. The entrance, however, of a sulphonic group in the molecule reducing its toxicity also reduces its therapeutic properties and this fact is in accordance with the general physiological inertia of the sulphonic acids

Sodium-allyl-thiocarbamino-p-stibanilate

Sodium allyl-thiocarbamino-p-stibanilate is another compound of probable therapeutic value which has been produced in my laboratory. The introduction of thio-urea may reduce the toxicity of the compound, just as it has been claimed in the case of the corresponding arsenic compound.

Glucose Derivatives

The therapeutic value of the glucose compounds of the organic aromatic antimonials, as compared with compounds

from which they are derived, is proportional to their antimony content and the same conclusion is arrived at on theoretical considerations. Their antimony contents are less than the corresponding aryl antimonials from which they are derived and therefore a bigger dose has to be administered to be of equal therapeutic value. The combination with glucose has therefore no advantage.

The antimony content of some of the aromatic antimonials is given below —

Sodium stibanilate	42 10 per cent
Urea strbamine	36 95
Chloro stibacetin (Stibosan)	33 30
Glueose sodiumi stibanilate	25 80
Gluense urea stibamine	23 80
N phenyl glyeine amide p stibinate of sodium	29 30
Neo stibosan	42 0

It may be stated that generally speaking the therapeutic value of the aromatic antimonials derived from p amino phenyl stibinic acid is proportional to their antimony content

As regards toxicity I have observed that the toxicity of pentavalent compounds obtained from p stibanilic acid is proportional to their antimony content. My observations are different from what I find stated in a recent book on kala azar, from which it will be seen that urea stibamine and its glucose derivatives are regarded to be equally toxic. The latter observation is rather significant as this would mean that the antimony content of the former is one and a half times less toxic than its glucose derivative.

It is a well known fact that the sodium salt of N phenyl p arsenic acid is a substance of practically no importance in 47–76 ~

the treatment of experimental infections such as those produced in laboratory animals by various species of trypanosomes, the spirochætes of relapsing fever and Tieponemapallidum On the other hand, N-phenyl-glycine-amide-parsenate of sodium, which is known under the trade name of truparsamide, is the most effective arsenical yet produced for the treatment of human sleeping sickness It has been stated that as Ehrlich's reduction theory proved fruitful in producing results of highly practical value, it does not represent the whole truth, for in tryparsamide, arsenic exists in the pentavalent form. But I hold that if a pentavalent organic arsenical, when introduced in the system, is more quickly converted into a compound containing the reactive - As = O than arseno-benzene compounds, then the former will be of more therapeutic value than the latter and on this the value of tryparsamide can be explained on Ehrlich's reduction theory.

Urea Stibamine

The most important and the last antimony compound that I shall now discuss is urea stibamine—lt is somewhat allied to tryparsamide

These compounds contain the group NH₂CO and, this, to my mind, is responsible for the therapeutic value of tryparsamide. That being the case, one would expect to have the same remarkably beneficial effects with antimony compounds containing this group in the treatment of diseases in which antimony is indicated, just as tryparsamide in the case of human trypanosomiasis.

This theoretical conclusion is borne out in practical experience. For to-day urea stibamine stands as the most pre-eminent compound of antimony in the treatment of kala-azar.

Taking into consideration the published kala-azar cases of different observers under different conditions and in

different places treated with the aramatic antimonials the mast extended trial has been given to ure a stibamine Observations upon the other aramatic antimonials are mostly limited to observations of single individuals

In a cambined series of 325 published cases which were treated by myself. Shortt. Greig. Kundu and others with this campound. 98 47 per cent of the cases were cured. One of the cases died of extreme nsthenia being admitted at the age af 65 in a maribund canditian. In 298 of these cases proof af cure was micrascapic and cultural eximinatians and disappearance af symptams and in 27 cases praaf of cure was clinical disappearance af the symptams and subsequent abservations of the cases. One case was resistant.

Regarding the value of N phenyl glycine amide p stibinate of sodium in the treatment af kala azar, I have published a series of eight cases in which it was successfully used but no comparison can be made nt present with urea stibamine as it has nat yet been given an extended trial in the treatment of the disease

It has been praved by the abservations of Shortt as well as those of myself and others that urea stibrimine does not manifest any deterioration or other changes either in physical and chemical characters or in therapeutic properties if kept in sealed ampaules under ordinary canditions. A more stable campound is undesirable as it will be less effective

I have already referred to my views af the reticulo endathelial system and I hold that individual cases will get beneficial results from the use of antimany campaunds pra partianal to the reaction of the reticula endathelial system. Twa things are necessary namely the development of the clasmatacytes and intraduction of an antimony compound with which they can cambine far the development of stibaxyl. Herein lies the value of the different antimonials and the superiority of urea stibamine over the other antimany.

compounds This also explains why, with the same antimony compound, one individual is cured much more quickly than another after its administration. It is the response of the cells of the reticulo-endothelial system to a particular drug that one should aim at in the treatment of the resistant cases.

Let me now briefly refer to the views of Voegtlin and his co-workers These observers have pointed out that arsenious oxide and its derivatives combine with substances containing a sulphydrile grouping and that the toxic action of the the simultaneous organic arsenoxides is depressed by injection of excess of sulphydrile compounds. Hopkins has shown that one such sulphydrile compound, reduced glutathione, plays an important part in the hydrolytic oxidation-reduction process of the living cell Voegtlin suggests that a combination of the arsenoxides with such groups and consequent suppression of this vital function may explain the toxic and curative actions of the arsenical derivatives, and that a formation by trypanosomes of the sulphydrile compound in excess of its vital need may be the basis of acquired resistance of trypanosomes The same probably takes place in the cases of Leishmania Investigations in these directions may lead to discovery of methods of preventing the development of antimony-resistant Leishmania

We have discussed the chemotherapy of antimony from a certain standpoint based chiefly upon the ideas of Ehrlich, Voegtlin and others. Other factors that have to be considered in this connection are the molecular weight of the compounds, their solubility, their dissociation in solution, their surface tension, the hydrogen-ion concentration of the tissue at which they act and various other points which, I am afraid, the time at my disposal will not permit me to discuss

I shall end here by quoting the remarks of Shortt and Sen which they made in 1925 about urea stibamine "We consider the value of urea stibamine has been established as the most efficient drug at present in use in the treatment of kala azar. This statement remains equally true to day. To this may be added the remarks of Dodds Price. I am of opinion that urea stibamine is a most valuable remedy in the resistant types of the disease and I strongly urge that it should be resorted to if after a few injections of sodium antimonyl tartrate a patient does not show marked improvement. I would only add that metallic antimony in a state of fine subdivision should be resorted to in those few cases which may be resistant to urea stibamine and which perhaps do not go beyond 3 per thousand or less.

It will be seen from what I stated that Frankel in his Agrentimettel is not justified in saying that changes in the molecular structure of antimony compounds do not bring about an increase in their therapeutic properties

[Proceedings of a meeting held at the Calcutta Medical Club on the 14th July, 1916]

Dr Upendranath Brahmachari read the following paper on "Colloids and other Drugs in the Treatment of Kalaazar," Sir Kailash Chandra Bose being in the chair

(ABSTRACT)

The remarkable bactericidal properties of electrical colloidal solutions of metallic silver led me to the use of electrargol in the treatment of kala-azar. So far, however, I have not been able to come to any definite conclusion about the therapeutic value of the drug in kala-azar Electro-mercural and electro-selenium have so far given only negative results.

All of us are aware of the remarkable effects of metallic antimony against the Leishmania and the Trypanosomes A colloidal preparation of this metal would therefore be an ideal drug in the treatment of kala-azar. Such a preparation would be comparatively free from toxic effects to the human organism and at the same time possess high bacterio-tropic properties against the Leishmania. The following cases fully justify these hopes.

Case No. I

Patient A, girl æt 16, was admitted for treatment of kala-azar Her spleen extended 6" below the costal arch and the splenic blood contained an unusually large number of L. D. bodies Her body weight was 4 stone. She was given 30 injections of colloidal antimony in doses of 002 grm to 003 grm on successive days. The result of the treatment was as follows —

- (1) Increase of body weight—1st 10 lb.
- (2) Temperature normal for nearly three months
- (3) Spleen gone down by nearly 3 inches

- (4) Disappearance of L D bodies from the splenic blood after 20 injections
- (5) R B C -4,100 000 W B C -3 200 Hb -40% on 29 2 16 (before treatment)

R B C -4,300 000 W B C -3 800 Hb -44% on 24 3 16 (at commencement of treatment)

RBC—4 300 000 WBC—7 400, Hb—48% on 14 4 16 RBC—4 600 000 WBC—7 000 Hb—58% on 9 6 16 (after treatment)

Case No 11

Patient B was admitted into my ward for treatment of kala azar on 16 2 16. He was cachectic and much emaciated at the time of admission. The spleen extended 3 below the costal arch and there was a large number of L. D. bodies in the splenic blood. He was at first treated with intramuscular and subsequently with intravenous injections of colloidal metallic antimony. Altogether 20 intravenous injections of the colloid (002 grm. each) were given. The result of the treatment was as follows.—

- (1) Temperature normal
- (2) Increase of body weight-6 lb
- (3) Disappearance of L D bodies from the splenic blood
- (4) R B C -2 600 000 W B C -4,800 Hb -40% on 17 2 16 (before treatment)

R B C —4 300 000 W B C —9 200 Hb —70% on 30 3 16 (after treatment)

Case No III

Patient T was admitted for treatment of kala azar The splenic blood showed presence of L D bodies. She had altogether 20 injections of the colloid (002 grm each) The result of treatment was as follows—

- (1) Temperature—normal
- (2) Spleen-diminished in size by two inches

(3) R B.C--1,600,000, W B C.--1,400, Hb --32% on 1.5 16 (before treatment).

R.B C -3,300,000, W B C -5,800, Hb -54% on 27 6.16 (after treatment)

It will thus be seen that colloidal metallic antimony produced remarkably beneficial effects on the patients

Another remarkable property possessed by the colloid is that it enables patients, who cannot bear treatment with tartar emetic or sodium antimonyl tartrate due to severe rigor, hyperpyrexia or severe vomiting setting in after the injections, bear treatment with these drugs if at first treated with three or four injections of the metallic colloid Probably the mild and non-toxic metallic colloid accustoms the patient to bear the subsequent administration of the soluble salts of antimony

Treatment of Kala-azai with Arsenic and Antimony Combined

In some cases of kala-azar, it has been found that after treatment with antimony, although, generally speaking, there was an all-round improvement, still the hæmoglobin value of the corpuscles could not be raised to the normal In such cases the combination of soamin with soluble salts of antimony was followed by very good results. In one case the hæmoglobin value was raised from 48 to 64 per cent after 8 injections of soamin, although it was at first almost impossible to raise the hæmoglobin value above 48 per cent with soluble salts of antimony alone. The effect of atoxyl on the treatment of kala-azar was described in a paper of mine published in the British Medical Journal some years ago. The paper was entitled Sporadic Kala-azar in Calcutta with notes of a case treated with Atoxyl

It may be stated here that colloidal metallic antimony has been prepared in a stable condition for the first time and been used for the first time by me in kala azar with the result I have just now mentioned

All of us are aware that the ointment of metallic antimony introduced by Sir Leonard Rogers produces markedly beneficial results in this disease. I have been able to prepare a non irritating ointment of tartar emetic and sodium emetic which also yielded similar results in this disease. In one case the ointment was rubbed for nearly a fortnight over the spleen the liver and the axilla on successive days as a result of which the spleen was diminished by 2 in size and the leucocytes rose from 2 600 to 5 600. The patient left hospital before the treatment was completed.

Conclusions

- (1) Colloidal metallic antimony has been obtained in a stable suspension for the first time. It produces remarkably beneficial effects in kala azar
- (2) It is perhaps the least toxic preparation of antimony and its use in kala azar is followed by tolerance towards the soluble salts of antimony sodium or potassium antimonyl tartrate in highly susceptible individuals
- (3) The combination of soamin with antimony rapidly improves the hæmoglobin value of blood in those cases of kala azar in which it is persistently low

It is not intended to refer in this paper to my experiences in the treatment of kala azar with metallic antimony sodium antimonyl tartrate and tartar emetic. These have been fully stated in my previous papers (Calcutta Medical Journal Oct and Nov. 1915 and the Indian Medical Gazette Dec. 1915 and January and May. 1916). I would however end by quoting what I have stated elsewhere namely that if future observations confirm the view that three or four

injections of metallic antimony are sufficient to bring about a complete and permanent cure of the disease, then we are in possession of a drug as powerful as quinine is for malaria, emetine for amoebic dysentery or salvarsan for syphilis—lts combination with formaldehyde will, perhaps, still more cut short the duration of the desease by the destruction of any antimony-fast parasites that may come into existence. To this I would add that the administration of soamin along with sodium or potassium emetic sometimes leads to a quicker improvement in the blood condition of the patient than is produced by the latter alone

NB—The method of preparation of a stable solution of colloidal metallic antimony will be described elsewhere

Discussion

Dr Rai Harmath Ghosh Bahadur remarked that he understood Dr Brahmachari to speak of the cure of kala-azar by only four or five injections of antimony. But in one of his cases he seemed to have mentioned that at least 20 injections were given to free the blood of the L D bodies. He did not thus realise fully the situation. In cases with hepatic enlargement, combination of emetine in one of his cases did accelerate the cure. He was very glad to find that antimony was bringing hopes about the discovery of the curative agent of the fell disease kala-azar.

Dr Rai Chunilal Bose Bahadur enquired how the colloidal antimony was prepared for Dr Brahmachari's use He had read in the Extra Pharmacopæia that of all colloidal preparations that of antimony was the most difficult to prepare Colloidal antimony had already been used in sleeping sickness. But here also the difficulty of obtaining it limited its use

Dr Rai Haridhan Dutt Bahadur remarked that the discourse given by Dr. Brahmachari that evening was very

convincing encouraging instructive and interesting. The cases which Dr Bralimachari had cited were certainly very convincing. But he wanted to know whether these successful cases were the only cases treated with colloidal antimony or these were among a group of several cases similarly treated but not with good results In private practice he had to deal with a few cases of kala azar Soamin and salvarsan treat ment both showed marvellous effects in some of his cases, but the effects did not last long and all died after some time So in the case of antimony treatment it was yet to be seen if the cures were permanent. Neo salvarsan was also used in three cases and he was struck with the marvellous improve ment but these too died after some time. He hoped that Dr. Brahmachari would be able to justify the remarks he made that evening after some time had elapsed. He hoped that as quinine was in malaria so Dr Brahmachari s colloidal atimony would turn out to be in kala azar

Dr U N Brahmachari in reply remarked that he was sorry to find that Dr Harmath Ghosh did not see the difference between colloidal antimony and metallic antimony in a state of fine subdivision. Three injections of metallic antimony could produce effects which twenty injections of salts of antimony could not do Regarding Dr H D Dutt's remarks he said that those were the only cases he had treated with colloidal antimony and in all of them he obtained good results. The future of colloidal antimony was great. It was non irritating While on the one hand it was the least organotropic on the other hand it was the most bacteriotropic. But still better preparations might be found hereafter previously read a paper about the treatment of kala azar by atoxyl With it the red and white corpuscles both increased but on the last day of the treatment when the patient was about to leave the hospital L D bodies were found in his blood Salvarsan and allied drugs have given some benefit but this did not last long. He had kept an open mind as his revered professor the late Dr Bomford used to advise.

Sir Kailash Chandra Bose in thanking the lecturer said that they all heard about the successful treatment of kala-azar by collodial antimony. It had a bright future. It would indeed be a glorious day for India, if an Indian could make further advances in the treatment of kala-azar

PREPARATION OF UREA ANTIMONYL TARTRATE A NEW COMPOUND

When excess of solid urea is added to a concentrated aqueous solution of hyper acid antimonyl tartrate and the mixture concentrated by heating on the water bath and then alcohol added to the mixture crops of prismatic crystals are obtained. These crystals are soluble in water and only very sparingly soluble in alcohol. They are best purified by being repeatedly washed with absolute alcohol. A solution of the salt gives a faintly acid reaction to litmus paper. On analysis the proportions of C. H. N and Sb present in the salt with the water of crystallization are as follows.—

{C=15 20% H=3 27% N=4 65% Sb=33 80% {Water of crystallization=12 25%

Calculated from CO(NH), (C,H; SbO O_c), 5H O which is assumed to be the chemical formula of the compound obtained

{C=15 00%, H=3 33% N=3 88% Sb=33 33% {Water of crystallization=12 50%

So far as I am aware there is no reference to this compound in the literature on compounds of urea and antimony

This salt is being used by me in the treatment of kala azar lts toxicity to lower animals seems to be rather low and experi ments are in progress to determine its toxic and curative doses

The solubility curve of the compound in water is shown in the accompanying chart

I am indebted to Mr Parimal Sen M Sc for helping me in the preparation of this compound and in working out its solubility curve

A CONTRIBUTION TO THE CHEMISTRY OF CERTAIN NEW AROMATIC ANTIMONIALS

The study of organic antimonials has not been so exhaustive as that of organic arsenicals In recent years some new organic pentavalent antimonials have been prepared and notable among these is urea stibamine discovered by Brahmachari, which has been found to be of great therapeutic value in the treatment of kala-azar. The reason why much less work has been done with organic antimonials than with arsenicals can be traced mainly to two important causes First of all, organic antimony compounds are very difficult to prepare and are with few exceptions not crystal-Secondly, most of them are unstable This instability limits the formation of various complex antimonials, which has been possible in the case of arsenic. This is especially the case with stibino-benzene compounds as compared with arseno-benzene compounds Generally speaking, case of arsenic, antimony, and bismuth this instability increases as the metallic character of the element becomes more and more pronounced Thus C-B1 link is less stable than C-Sb link and C-Sb link is less stable than C-As link

The great difficulty involved in the preparation of aryl antimonials is really a barrier against extensive investigations on this type of compound. This difficulty becomes still greater, as minute impurities and slight variations of physical influences affect the stability of the compounds to a considerable extent, thereby bringing about marked changes in their toxicity and therapeutic properties

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In the Indian Journal of Medical Research, the Indian Journal of Medicine and the Calcutta Medical Journal a series of new organic antimonials were described some time ago by Brahmachari and some of these compounds have been shown to be of great therapeutic value. Another series of new aromatic antimonials have since been investigated in the Brahmachari Research Institute and the following form the first series of such compounds.

- l Disodium p amino phenyl stibinate N methylene sul phonate
 - Urca p amino phenyl stibinate N methylene sulphon ateof sodium
 - 3 Disodium p stibinilate N methylene-sulphinate
- 4 Urea p amino phenyl stibinate N methylene sulphin ate of sodium
 - 5 p Acetyl amino plienyl stibinate of urca
- 6 1 Acctamino 2 azobenzene 4 4 distibinate of sodium
- 7 p l-lydroxy plienyl stibinate of urea

Some of these compounds as will be seen from their percentage composition given below, exhibit strong polymerisation whereby three molecules associate together giving rise to more complex molecules

EXPERIMENTAL

(1) Disodium p amino plienyl stibinate N methylene sul

3(4 SO_aNa CH₂ NH C₁H₄ SbO₂)IH ONa

The starting material in the preparation of this compound is stibanilic acid which has been prepared by Bart's reaction. Stibanilic acid is neutralised with solution of sodium hydroxide and the sodium salt precipitated by absolute alcohol. The precipitate is then thoroughly washed with absolute alcohol till filtrate is free from alkali. It is next dried in a vacuum dessicator.

Sodium stibanilate is dissolved in water and then formal-dehyde solution and NaHSO, dissolved in water are added to it successively in a flask. The mixture is next heated on a water bath and filtered. The filtrate is treated with excess of alcohol when a bulky precipitate is produced which is washed with alcohol and dried in a porous plate in a vacuum dessicator.

The product is a light coloured powder—easily soluble in water to a perfectly clear solution which gives a faintly acid reaction

Composition —

Dried material corresponds to the formula

$$(SO_7Na\ CH_2\ NH\ C_tH_1.SbO)_1O_2(OH)(ONa)$$

= $C_{21}H_{22}O_{16}N_1S_7Sb_7Na_1$

Calculated for
$$C_{21}H_{22}O_{16}N_5S_5Sb_1Na_1 - Sb = 322\%$$
, $S = 87\%$, $N = 38\%$

Found-

$$Sb = 325\%$$
, $S = 85\%$, $N = 40\%$

(2) Urea p-amino-phenyl-stibinate-N-methylene sulphonate of sodium.

3(4-SO₃Na CH₂NH C₁H₁SbO₂)H OH₁N CO NH₂

The starting material in this preparation is urea stibamine. Urea stibamine is dissolved in a small quantity of water to which formaldehyde solution and NaHSO₀ dissolved in little quantity of water are added in succession. A bulky precipitate is formed on adding the constituents. The mixture is warmed on water bath. It is next filtered and the filtrate precipitated by alcohol. The precipitate is washed with absolute alcohol and then dried over a porous plate in a vacuum dessicator.

The product is a light coloured powder, easily soluble in water and gives a neutral reaction to litmus paper

Composition —

Dried material corresponds to the formula

SO, Na CH NH C, H, SbO), O (OH) (ONH, CO NH)

= C₂₂H₂O₃ N S₃Sb₃N₃s

Calculated for C-H-O₁ N₃S Sb₃Na₃ Sb=311% S=829% N=60%

Found

Sb=317% S=81% N=63%
(3) Disodium p stibanilate N methylene sulphinate
3(4 SO,Na CH NH C,H,SbO)H ONa

Stibanilic acid is treated with NaOH solution and the sodium salt next precipitated by adding absolute alcohol. The precipitate is washed with alcohol to remove the free alkali. The dried sodium salt is then dissolved in a little water and the solution thus obtained treated with sodium formaldehyde sulphoxylate dissolved in little water. A bulky precipitate appears and the whole mixture is warmed on a water bath when a clear solution is obtained with a small quantity of insoluble impurity. The solution after filtra tion is slightly concentrated and then precipitated by absolute alcohol. The precipitate is next filtered, and dried over a porous plate in a vacuum dessicator.

The product is a light coloured powder readily soluble in water to a perfectly clear solution which is neutral to litmus

Composition -

Dried material corresponds to the formula (SO Na CH₂ NH C_0 H₄ SbO)₄O (OH)(ONa) = C_1 H₂O₁₅N₅S₅Sb₅Na₄
Calculated for C_1 H O₁₅N₅S₅Sb₅Na₁
Sb=337% S=90% N=4%

Found

Sb=335% S=94%, N=42%

(4) Urea p-amino-phenyl stibinate-N-methylene sulphinate of sodium

3(4-SO₂Na CH₂ NH C₆H₄ SbO₂)H ONH₅ CO NH₂

Urea stibamine is dissolved in water to which a solution of sodium formaldehyde sulphoxylate is added. A bulky precipitate appears and the whole mixture is well shaken. The mixture is next warmed on a water-bath. A clear solution with a slight sediment at the bottom is obtained which is next filtered. The clear filtrate after concentration is precipitated in cold by absolute alcohol. The precipitate is washed with alcohol, and dried over a porous plate in a vacuum dessicator.

The product is a light coloured powder, readily soluble in water to a perfectly clear reddish solution which is faintly acid to litmus

Composition —

Dried material corresponds to the formula (SO₂Na CH₂ NH C₆H₄ SbO)₇O₂(OH)(ONH CO NH₂) = C₂₂H₂₂O₁₁N₆S₃Sb₃Na₃

Calculated for C22H27O14N5S2Sb-Na-

$$Sb = 32.4\%$$
, $S = 3.65\%$, $N = 6.3\%$

Found

$$Sb = 320\%$$
, $S = 34\%$, $N = 60\%$

(5) p-Acetyl-amino-phenyl-stibinate of urea 3(4-CH₂CO NH C₆H₂ SbO₂)H ONH₂ CO NH₂

The starting material in this preparation is p-acetyl-aminophenyl stibinic acid which is obtained from the corresponding acetyl-phenylene-diamine. The acid is thoroughly washed and the pasty mass is obtained in a semi-dry state by pressing over porous plate. The moist acid is treated with a little urea and then well mixed. The mixture is heated in boiling water when a reddish solution is obtained. A little more water may be added, if necessary, to obtain a clear solution and then warmed. The solution is next

filtered through fluted filter paper and the filtrate precipitated by absolute alcohol The precipitate is well washed with the same and dried over a porous plate in a vacuum dessicator

The product is a yellowish powder and dissolves in water to a clear solution which is faintly acid

Composition —

Dried material corresponds to the formula (CH, CO NH C_cH_s SbO)₂O₂ (OH)(O NH₂ CO NH) = $C_{25}H_{\infty}O_{11}N$ Sb₃

Calculated for C₂H₂O₁₁N Sb₃ N=748% Sb=384%

Found

N=79%, Sb=380%

(6) 1 Acetamino 2 azobenzene (4 4") distribinate of sodium CH, CO NH C₂H, N N C₂H₃(SbO₂HNa)₂
= C₁H₁·C₂N₂Sb Na

The starting materials in the preparation of this compound are acetyl stibanilic acid and stibanilic acid. The former is obtained from acetyl p phenylene diamine and the latter by its hydrolysis with alkali. The stibanilic acid is partially dried on a porous plate and suspended in a small quantity of water. The mixture is cooled and treated with excess of H₂SO₄ when a clear solution is obtained Acetyl stibanilic acid dried similarly is weighed and then dissolved in excess of alkalı The former acid solution is then gradually treated with NaNO solution till it gives a blue coloration with the starch todide paper. The alkaline solution of the acetyl stibanilic acid is also cooled in ice and then gradually added to the diazotised solution. It is then filtered after allowing the little quantity of froth to escape. The sodium salt is then precipitated from the concentrated solution by absolute alcohol-dried over a porous plate in a vacuum dessicator

The product is a brown powder which dissolves in water giving a clear red solution with neutral reaction

Composition —

Calculated for $C_{11}H_{10}O_7N_0Sb_2Na_2$ N = 6.76%, Sb = 38.6%Found N = 7.0%, Sb = 38.1%

(7) p-Hydroxy-phenyl-stibinate of urea 4-OH C₆H₁SbO₅H NH₃ CO.NH₂

p-Stibanilic acid which is obtained from acetyl-p-phenylene-diamine is made into a thick paste with water and an excess of H₂SO₁ added, the mixture being cooled. A solution is produced in this way which is well stirred while NaNO₂ solution is gradually added till it imparts a blue colour to starch-iodide paper immediately. The mass is next dissolved in alkali after gentle warming to liberate all nitrogen and filtered. The filtrate is reprecipitated with acetic acid. The mixture is filtered and well washed with water. The hydroxy-phenyl-stibinic acid thus obtained, which can also be directly obtained from p-amino phenol by applying Bart's reaction, is then well mixed with little excess of urea and heated on water bath when a clear red solution is obtained. It is then filtered and precipitated by acetone and dried in vacuo over a porous plate.

The product is a yellow powder readily dissolving in water to a perfectly clear solution which is faintly acid to litmus

Composition — Calculated for $C_7H_{11}O_5N_2Sb$ N=87%, Sb=37%. Found N=9%, Sb=376%.

The therapeutic value, if any, of these compounds will be reported later on

References

⁽¹⁾ Indian Journal of Medical Research, Vol. X, No. 2, Oct., 1922, Vol. XI, No. 1, July, 1923. Vol. XI, No. 2, Oct., 1923. Vol. XI, No. 4, April 1924. Vol. XII, No. 1, July 1924, Vol. XII, No. 2, Oct., 1924, Vol. XII, No. 4, April, 1925, Vol. XIII, No. 1, July, 1925, Vol. XIII, No. 3, January, 1926.

⁽²⁾ Indian Journal of Medicine, June, 1926, Sep., 1926

⁽³⁾ Calcutta Medical Journal, June, 1926, Aug., 1926

SYNTHESIS OF A FEW ANTIMONIALS OF THERAPEUTIC INTEREST

This paper contains an necount of some organo metallic antimonials which have been synthesised with the same object in view as in the case of the compounds described in a previous paper contributed to this Journal (Vol XXV 1929 No 1) They are amorphous and ex tremely difficult to purify The chemical operations involved in their preparation are given below. As regards their toxicity we have noticed that in these as in the previous compounds the general rule holds or, introduction of sulphoxyl groups lowers the toxicity to n considerable extent with in decrease in the therapeutic value. The nature of the basic portion also affects to some extent the stability and the toxicity of the compounds, via uren or diethylamine salt is sometimes more stable and less toxic than the corresponding sodium salt Our object in the preparation of the following compounds is to study these latter effects as well. The compounds are not very stable though their solutions do not decompose on standing in air for 24 hours. The following is a list of the compounds investigated by us in this paper -

- Sodium salt of phenyl glycine amide 4 stibinic acid
- 2 Urea salt of the same
- 3 Diethyl amine salt of the same
- 4 Carbamino p stibanilate of sodium
- 5 Carbamino p stibanilate of urea
- 6 Carbamino p stibanilate of diethyl amine

It will be seen that all the above compounds undergo polymerization (see below)

EXPERIMENTAL

(1) Sodium phenyl-glycine-amide-4-stibinate

p-Stibanilic acid is dissolved in the requisite quantity of NaOH solution and the concentrated solution of sodium p-stibanilate is added gradually to an excess of absolute alcohol, when a precipitate of sodium stibanilate is produced, which is next filtered and washed with absolute alcohol and then died 5 grms of sodium stibanilate are then dissolved in methyl alcohol and treated with chloracetic ester and the whole refluxed for several hours After the reaction is complete, the methyl alcohol is distilled off and the rest acidified with dilute HCl The precipitate thus obtained is filtered and washed with water and then treated with concentrated ammonia After some time, the solution is filtered and the filtrate is reprecipitated by acetic acid, when the glycine amide derivative is obtained which is next washed with distilled water The precipitate is then dissolved in dilute NaOH, filtered and the filtrate precipitated by adding absolute alcohol The precipitate is then repeatedly washed with absolute alcohol and dried in a vacuum desiccator

It is an almost white powder, very easily soluble in water to a perfectly clear solution, which gives neutral reaction with litmus. On warming with dilute alkali it gives out ammonia. The compound prepared according to the above process has been called X_{10} , a paper on the thera-

peutics of which has been published in the Transactions of the Royal Society of Tropical Medicine and Hygiene The method of preparation of the compound described here is better than the one originally described by Brahmachari in the Indian Journal of Medical Research 1922

Composition -

Dried material corresponds to the formula

(NH CO CH, NH C,H SbO), O₂(OH₁(ON_t)

=C₂,H₁,O N₄Sb₄Na

Found Sb=38 40 ' N=8 72 o Calculated for C . H . O. N Sb Na Sb=38 17 N=8 90

This compound is the polymerized antimony analogue of tryparsamide

(2) Phenyl glycine amide 4 stibinate of urea (NH CO CH NH C,H SbO),O,(OH)(ONH, CO NH,)

Phenyl glycine amide 4 stibinic acid as obtained in the previous experiment is made into a paste with little water and then well mixed with an excess of urea. The whole is then warmed for some time when the acid gradually dissolves to a reddish solution yielding a urea salt. The solution is then filtered through a Buchner funnel and the clear filtrate is precipitated by acetone. The precipitate thus obtained is dried in a vacuum desiccator after washing well with absolute alcohol.

The product is a light coloured powder easily dissolving in water to a perfectly clear solution which is neutral to litmus

Composition ---

Dried material corresponds to the formula

(NH CO CH, NH C₄H SbO) O (OH)(ONH CO NH)

=C₂₃H, O₃₁N Sb₃

Found Sb=36 589' N=11 55 /

Calculated for C25H 3O N Sb Sb=36 69 / N=11 41 /6

(3) Phenyl-glycine-amide-4-stibinate of diethyl amine (NH₂CO CH₂ NH C₆H₁ SbO)₃O₂(OH) OH₂N(C₂H₅)₂

The starting material in the preparation of this compound is the same as in the previous cases. This is well mixed with a small quantity of water, and to the mixture a 30 per cent solution of diethylamine in water is gradually added, shaking it very well at the same time. Almost a clear concentrated solution is thus obtained, which is filtered and the reddish filtrate is poured drop by drop into 5 times its volume of absolute alcohol. A voluminous precipitate is produced, which is allowed to settle down for some time and then filtered. The precipitate is washed well with absolute alcohol and then dried in a vacuum desiccator.

It is a light grey powder easily dissolving in water to a clear solution which is neutral to litmus

Composition ---

Dried material corresponds to the formula

$$(NH_2CO CH_2 NH C_6H_4 SbO)_3O_2(OH) OH_2N(C_2H_5)_2$$

= $C_{28}H_{40}O_{10}N_7 Sb_3$

Found Sb=36 42%, N=9 71% Calculated for C₂₈H₄₀O₁₀N₇Sb₃ Sb=36 21%, N=9 85%

(4) Sodium carbamino-p-stibanilate

The starting material in the preparation of this compound is sodium p-stibanilate, produced by neutralising p-stibanilic acid with NaOH solution, the acid itself being obtained by

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hydrolysing acetyl p stibanilic acid and which is a product of Bart's reaction applied to acetyl p phenylene diamine Five grms of sodium stibanilate thus obtained are dissolved at low temperature in glacial acetic acid. To this well cooled mixture are gradually added about 4 grms of potassium cva nate and the mixture well stirred till a clear solution is obtained. The solution is then allowed to remain in this state for several hours. The mixture is then diluted with water and well stirred Concentrated HCl is then gradually added which dissolves the unreacted p stibanilic acid and precipi tates the carbamino derivative as a voluminous mass which is then filtered and washed with water. The wet precipitate is then dissolved in the requisite quantity of dilute NaOH solution and the reddish solution thus obtained is filtered The filtrate is precipitated by absolute alcohol and the preci pitate washed with the same and then dried in a vacuum desiccator

The product is almost a white powder readily dissolving in water to a clear solution which is neutral to litmus

Composition -

Dried material corresponds to the formula (NH₂ CO NH C₄H₄ SbO) O₂ OH ONa

=C₁H₂₂O₁₀N₆Sb₃N₈

Found Sb=39 62% N=9 29%
Calculated for C₂₁H₂₂O N₆Sb₃Na Sb=39 95 / N=9 32 /₆

(5) Carbamino p stibanilate of urea
(NH, CO NH C H, SbO), O OH ONH, CO NH,

Carbamino p stibanilic acid as obtained in the previous case is made into a paste with a little water and then well mixed with a slight excess of urea. The mixture is then warmed on a water bath when the acid gradually dissolves to a clear solution. The solution is next filtered and the filtrate precipitated by acetone.

It is a light grey powder which dissolves easily in water giving a neutral solution.

Composition —

Dried material corresponds to the formula

$$(NH_2 CO NH C_rH_4 SbO)_7 O_2 OH.ONH_7 CO NH_2$$

= $C_{22}H_2 \cdot O_{11}N_6Sb_7$

Found Sb = 3850%, N = 1185%.

Calculated for $C_{22}H_{27}O_{11}N_8Sb_7$ Sb=38 34%, N=11 92%

(6) Carbamino-p-stibanilate of diethyl amine (NH₂ CO NH C₁H₄ SbO)₃ O₂ OH ONH₂ (C₂H₄)₂

As in the previous experiment a paste is made by mixing carbamino-p-stibanilic acid with little water to which is then gradually added a 35 per cent solution of diethyl-amine in water. The precipitate gradually dissolves, giving a clear solution which is filtered, and the filtrate reprecipitated by acetone

It is a pale greyish powder which dissolves readily in water.

Composition —

Dried material corresponds to the formula:

$$(NH_2 CO NH C_6H_4 SbO)_3 O_2 OH ONH_2 (C_2H_5)_2$$

= $C_{25}H_{34}O_{10}N_7Sb_3$

Found Sb=37.62%, N=10.31%

Calculated for C_2 , $H_{34}O_{10}N_7Sb_3$ Sb=37.81%, N=10.29%

References

- 1 Journal and Proceedings of the Asiatic Society of Bengal (New Series), Vol XXV, No. 1, 1929
- 2 Transactions of the Royal Society of Tropical Medicine and Hygiene, Vol XXIII, No 6, pp 617-622, April, 1930,

SYNTHESIS OF SODIUM N-PHENYL-GLYCINE AMIDE 4 STIBINATE (ANTIMONY ANALOGUE OF TRYPARSAMIDE)

This paper gives the method of preparation of one of a series of new antimonials of therapeutic interest which were under publication in the Jaurnal and Proceedings of the Asiatic Society of Bengal (Vide page 390)

The above compound has been found to be of thera peutic value in the treatment of kala azar and a series of cases has been published in the Transactions of the Rayal Society of Trapical Medicine and Hygicac Vol XXIII No 6 pp 617 622 April 1930 A paper containing a further series of cases of kala azar treated with this compound is under preparation

[NB—Another method of preparation of the above compound by Brahmachari was described in the Indian Journal of Medical Research October 1922—Ed]

THE INTENSIVE ANTIMONIAL TREAT-MENT OF KALA-AZAR

PART I

In the Indian Journal of Medical Research (1925), Brahmachari described a series of cases of kala-azar treated by an intensive course with urea stibamine, which consisted of injections given daily or on alternate days or of multiple injections given on the same day during a course ranging from thirty-six hours to seven days

Since the publication of the above paper one hundred and twenty-five more cases in which the intensive treatment by daily administration of the above drug produced similarly satisfactory results, consisting of lessening of the period of treatment which extended from seven to ten days and without any untoward symptoms due to the short-interval injections, have been recorded.

The method of treatment by daily injections was subsequently followed by Napier and Mullick with another antimonial, neo-stibosan, and they reported that they likewise had got satisfactory results in their cases

It may at once be mentioned here that cases of kala-azar vary most markedly in their response to specific treatment, as has been pointed out in this Journal and elsewhere. This variability is perhaps more noticeable in kala-azar than in any other protozoal disease. It is very important to remember this, as otherwise medical men may be tempted to believe that a case that has had a week's or ten days' treatment with a pentavalent aromatic anti-

monial is cured or one may be disappointed if one finds that a particular case is not responding as quickly as one would expect from a study of the reported cases in which cure has been recorded by a short course of treatment

As illustrations of what may be regarded as easily cured cases of the disease the notes of a few cases that recently came under our observation are appended here —

- (1) Patient act 6, was admitted into the wards of the Chittaranjan Hospital with history of fever with double rise of temperature for 9 months Spleen extended 31 in below the costal arch L. D bodies were found on spleen puncture Blood examination showed R B C -1 200 000 WBC-2 872 and Hb-40 per cent Patient was given daily intravenous injections of urea stibamine for 7 days in doses of (1) 075g (2) 1g (3) 15g (4) 2g (5) 2g (6) 2g the total amount being 925 gramme after which the treatment was stopped. Patient was kept under observation for one month and a half. At the time of discharge spleen could not be felt below the costal arch no L D bodies could be found on spleen puncture and W B C count was 8 000 per c mm Patient was free from fever during the period of observation after the course of treatment
- (u) Patient S was treated in the Health Association Ward No. 4 Calcutta Municipality with history of fever with daily double rise of temperature for 9 months. Spleen extended 41 in below the costal margin. Patient was given daily intravenous injection of urea stibamine for 6 days in doses of (1) 025g. (2) 025g. (3) 025g. (4) 025g. (5) 05g. (6) 05g the total amount being 2 gramme after which the treatment was stopped. Patient was kept under observation for three months. At the time of discharge spleen could not be felt below the costal margin. Patient was free from fever during the whole period of observation after the course of treatment.

- Hospital with history of fever with daily double rise of temperature for 9 months. Spleen extended 4 in. below the costal arch and liver just palpable. L. D. bodies were found on spleen puncture. Blood examination showed R.B. C.—2,800,000, W.B.C.—3,120 and Hb.—45 per cent Patient was given daily intravenous injection of urea stibamine for 7 days in doses of (1) '025g, (2) '05g, (3) '1g., (4) 1g., (5) 1g, (6) '1g, (7) '1g., the total amount being 575 gramme, after which treatment was stopped. Patient was kept under observation for 33 days. At the time of discharge, spleen could not be felt below the costal arch, no L. D. bodies could be found on spleen puncture and W.B.C. were 6,552 per c. mm. Patient was free from fever during observation.
- (10) Patient P, æt. 29, was admitted in the Chittaranjan Hospital with history of fever with daily double rise of temperature for one year and two months Spleen extended $3\frac{1}{2}$ in below the costal arch Blood examination showed R B C —3,100,000, W B C —3,432 and Hb —45 per cent L D bodies were found on spleen puncture Patient was given daily intravenous injections of urea stibamine for 8 days in doses of (1) 05g, (2) 075, (3) 1g, (4) 1g, (5) 1g., (6) 15g, (7) 2g, (8) 2g, the total amount being 975 gramme, after which the treatment was stopped Patient was kept under observation for 27 days At the time of discharge, spleen could not be felt below the costal arch, no L D bodies could be found on spleen puncture and W.B C were 6,573 per c mm Patient was free from fever during observation
 - (v) Patient K, æt 30, was admitted in the Chittaranjan Hospital with history of fever with daily double rise of temperature for 7 months He had previously 12 injections of neo-stibosan Spleen extended $4\frac{1}{2}$ in below the costal margin at the time of admission Blood examination

showed R B C —3 600 000 W B C —2 184 and Hb —65 per cent L D bodies were found on spleen puncture Patient was given daily intravenous injections of urea stiba mine for 9 days in doses of (1) 025g (2) 05g (3) 05g (4) 1g (5) 15g (6) 2g (7) 15g (8) 2g (9) 2g the total quantity being I 125 grammes, after which treatment was stopped Patient was under observation for two months At the time of discharge spleen could not be felt below the costal arch no L D bodies could be found on spleen punc ture and W B C was 5,800 per c mm

(vi) Patient N set 7 was admitted in the Chittaranjan Hospital with history of fever with daily double rise of tem perature for 8 months. Both spleen and liver extended 2 in below the costal arch. L D bodies were found on spleen puncture. Blood examination showed R B C —1 720 000 W B C —2 500 and Hb —35 per cent. Patient was given daily intravenous injections of urea stibamine for 9 days in doses of (1) 025g (2) 05g (3) 075g (4) 1g (5) 1g (6) 1g (7) 1g (8) 1g (9) 1g the total amount being 75 gramme after which the treatment was stopped. The patient was kept under observation for two months. At the time of discharge spleen could not be felt below the costal arch no L D bodies could be found on spleen puncture and W B C count was 6 056 per c mm. There was no fever during the period of observation.

On the other hand there are cases in which a prolonged course of treatment with any antimonial aromatic or otherwise only slowly influences the course of the disease. It is to these cases that attention of research workers should be directed at the present day especially in view of the fact that these cases are frequently not properly recorded by those who advocate a particular antimonial preparation which failed to act quickly in them

It has been frequently held that early cases are easily amenable to treatment—a view originally expressed by

Brahmachari in a paper published in the Indian Journal of Medical Research in 1924 under the title of "Value of Urea Stibamine in the Treatment of Early Kala-azar" Further experience has, however, shown that while this may be so in a large number of patients, there are certain cases which are refractory from the very beginning.

Further, in an early case, one sometimes observes that a few weeks' delay in commencing specific treatment helps in bringing about a cure more quickly than immediately starting the treatment.

This brings one to the question of production of antibodies in the treatment of kala-azar. Just as in the case of treatment of syphilis with an arsenical, or of malaria with quinine, little work has been done in the production of antibodies in the treatment of kala-azar with an antimonial.

The object of this paper is to indicate that while the intensive treatment of kala-azar with urea stibamine cuts short the course of treatment of the disease as quickly as any other antimonial that has now been put on the market, it would be misleading to assert that such brilliant results are to be met with in all cases, whatever that antimonial preparation may be

REMARKS

The mechanism of response of leishmania to an antimonial preparation is a very complicated one. While an aromatic pentavalent antimonial, such as urea stibamine, brings about sterilization of an infected individual in a much shorter time than tartar emetic, it must, at the same time, be admitted that, with any such compound, the time required for sterilization is variable in the case of different individuals. What is the mechanism of this variability. This constitutes an important line of research, and will be referred to in a subsequent paper.

References

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B hm ch r U N & M ty B B 11925] Chemoth app of Antmon I Compound in K la atar Infection Pat XIV—Observat in on a series of ca of Kalvaza tr t d with Urea Stb mine during a coule of 32 hou to 7 d y Ib d Apr 1 1925

THE INTENSIVE ANTIMONIAL TREATMENT OF KALA-AZAR

PART II

This paper is a continuation of a previous paper by Brahmachari and a co-worker published in this Journal It gives a collective series of 31 (1931) cases of kala-azar treated with urea stibamine by the intensive method, in addition to those previously recorded cases were treated in the Tropical Diseases Ward of the Carmichael Medical College Hospitals, the Kala-azar Ward of the Chittaranjan Hospital, the Campbell Hospital, Calcutta, and in the Out-Patient Department of the Chittaranjan Hospital, Calcutta It was intended to test the value of the drug by the intensive method independently and give a collective report. The present paper gives a report of cases treated up to the time of writing Further observations are still in progress, and will be reported in another series

(a) Notes on Cases treated in the Tropical Diseases Ward of the Caimichael Medical College Hospitals, and in the Kala-azar Ward of the Chittaranjan Hospital, Calcutta

CASE No 1—J N D, æt 19, was admitted on 22-9-32 History of irregular fever for about nine months. Patient anæmic, liver extended $1\frac{1}{2}$ in and spleen 3 in below the costal arch. Patient developed pleurisy in hospital before he was treated for kala-azar. Spleen puncture L D, bodies present. After the patient was cured of

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pleurisy, he was treated with urea stibamine injected intravenously on successive days from 4 10 32. The following were the doses given (1) 0 05 g (2) 0 05 g (3) 0 1 g (4) 0 1 g, (5) 0 1 g (6) 0 15 g, (7) 0 15 g (8) 0 15 g (9) 0 15 g and (10) 0 2 g. Altogether ten injections were given with a total amount of 1 2 grms. The temperature came down to normal after the first injection and remained so up to the day of discharge from hospital. No reactions were observed during treatment. At the time of discharge spleen could not be felt below the costal margin and no L. D. bodies were found on spleen puncture. Patient remained in hospital for two months after the completion of treatment.

CASE No 2 -N art 36 was admitted on 3 9 32 History of irregular fever for about a year with double rise of temperature pyorrhoea alveolaris and bleeding from gums present Patient anæmic Liver extended 1 in and spleen 6 in below the costal arch in the mid clavicular line Spleen puncture L D bodies present Patient was treated with urea stibamine injected intravenously on successive days from 7 9 32 The following were the doses (1) 0 05 g (2) 0 1g (3) 0 1 g (4) 0 1 g (5) 0 15 g (6) 0 15 g (7) 0 15 g (8) 0 15 g (9) 0 15 g (10) 0 2 g and (11) 0 2 g Al together eleven injections were given with a total amount of 1.5 grms The temperature came down to normal after the second injection and remained so up to the day of discharge from hospital No reactions were observed during treat ment At the time of discharge spleen extended 2 in below the costal margin and no L D bodies were found on spleen puncture The patient remained in hospital for two months after the completion of treatment

Case No 3—A C P act 20 was admitted on 31 8 32 History of irregular fever for 1 year About three months ago patient had general anasarca Liver was tender extending 1 in and spleen 6 in below

Patient was treated with urea stibamine injected intravenously on successive days from 7-8-32. The following were the doses (1) 0 05 g, (2) 0 l g, (3) 0 l g, (4) 0 l g, (5) 0 l 5 g, (6) 0 l 5 g, (7) 0 l 5 g, (8) 0 l 5 g and (9) 0 2 g Altogether nine injections were given with a total amount of 1 l 5 grms. The temperature came down to normal after the second injection and remained so up to the day of discharge from hospital. At the time of discharge spleen could just be felt below the costal margin, but no L. D bodies were found on spleen puncture. Patient's general condition very much improved. Patient remained in hospital for two months after the completion of treatment.

CASE No 4.—A M, æt 14, was admitted on 22-8-32 History of irregular fever Patient was ill-nourished At the time of admission, ascites and cedema of the feet were present with old scars over the abdomen in the splenic area Liver was hard, its margin extending \frac{1}{2} in below the costal margin and spleen was very hard, reaching the level of the umbilicus and \frac{1}{2} in. from the middle line to the right Spleen puncture L. D bodies present. There was presence of albumin in the urine. Patient was treated with urea stibamine injected intravenously on successive days from 28-8-32 During treatment ascites and ædema disappeared The following were the doses (1) 0 05 g, (2) 0 05 g, (3) 0 05 g., (4) 0 1 g, (5) 0 1 g, (6) 0 1 g, (7) 0 1 g, (8) 0.15 g, (9) 0.15 g, (10) 0.1 g, (11) 0.1 g and (12) 0 15 g Altogether twelve injections were given, with a total amount of 1 15 grms. The temperature came down to normal after the fourth injection and remained so up to the day of discharge from hospital No reactions were observed during treatment At the time of discharge, spleen extended one inch below the costal margin and no L D bodies were found on spleen puncture Patient remained in hospital for two months after the completion of treatment

CASE NO 5-JAM set 22 was admitted on 2932 History of irregular fever and general anasarca At the time of admission patient was aniemic with ædema of the face and limbs Liver was hard and extended 2 in and spleen also very hard extended 7 in below the costal margin in mid clasicular line and 4 in to the right of the middle line. Spleen puncture L D bodies present Patient was treated with urea stibamine injected intravenously on successive days from 7932 The following were the doses (1) 000 g (2) 0 lg (3) 0 lg (4) 0 lg (5) 0 lg (6) 0 lg (7) 0 15 g (8) 0 15 g (8) 0 15 g (9) 0 15 g (10)0 15 g and (11) 0 2 g Altogether eleven injections with a total amount of 1 35 grms were given. The temperature came down to normal after the first injection and remained so up to the day of discharge from hospital No reactions were observed during treatment. At the time of discharge spleen could be felt about 2 in below the costal margin but no L D bodies were found on spleen puncture Patient remained in hospital for one month after the comple tion of treatment

Case No 6—G N K ret 14, was admitted on 29 8 32 History of irregular fever for about three years Spleen extended 91 in and liver enlarged to the finger's breadth below the costal arch Spleen puncture L D bodies present Patient was treated with urea stibamine injected intravenously on successive days from 6 9 32. The following were the doses (1) 0.05 g (2) 0.05 g (3) 0.05 g (4) 0.05 g (5) 0.1 g (6) 0.1 g (7) 0.1 g (8) 0.15 g (9) 0.15 g and (10) 0.15 g. Altogether ten injections with a total amount of 0.95 g were given. The temperature came down to normal after the first injection and remained so up to the day of discharge from hospital. No reactions were observed during treatment. At the time of discharge spleen was 2 in

below the costal margin and no L D bodies were found on spleen puncture. Patient remained in hospital for one month after the completion of treatment

CASE No 7 — T G, æt. 20, was admitted on 30-6-32. History of irregular fever with double rise of temperature during the first seven months. Spleen extended 51 in below the costal margin and liver not enlarged Patient moderately anæmic Spleen puncture L. D bodies present Patient was treated with urea stibamine, injected intravenously on successive days from 6-9-32 The following were the doses. (1) 0.15 g, (2) 0 15 g, (3) 0.15 g, (4) 0 15 g, (5) 0.15 g, (6) 0 15 g, (7) 0 2 g, (8) 0 2 g, (9) 0 2 g, and (10) 0.2 g. Altogether cen injections were given with a total amount of 17 grms The temperature came down to normal after the second injection and remained so up to the day of discharge from hospital No reactions were observed during treatment. At the time of discharge, spleen was 1 in below the costal margin, but no L D bodies were found on spleen puncture Patient remained in hospital for one month after the completion of treatment

CASE No 8.—B. K D., at 25, was admitted on 26-9-32. History of continued fever for five months and of double rise of temperature for one month. Spleen extended 5½ in below the costal margin and liver was just palpable. Spleen puncture L. D bodies present Patient was treated with urea stibamine, injected intravenously on successive days from 28-9-32 During this period, he had an attack of influenza and the injections were stopped for the time being. After the subsidence of bronchitis, injections of urea stibamine were continued from 21-10-32 to 24-10-32. The following were the doses. (1) 0.025 g, (2) 0.05 g., (3) 0.1 g, (4) 0.1 g, (5) 0.1 g., (6) 0.15 g., (7) 0.1 g, (8) 0.15 g., (9) 0.2 g and (10) 0.2 g Altogether ten injections with a total amount of 1.175 grms were given. The temperature came down to normal after the third injection and

remained so up to the day of discharge from hospital. No reactions were observed during treatment. At the time of discharge spleen was 2 in below the costal margin and no L. D. bodies were found on spleen puncture. Patient remained in hospital for one month after the completion of treatment.

CASE NO 9 -B P R, set 15, was admitted on 20 9 32 History of fever of an intermittent type occurring irregularly for about one year with bleeding from the gums Spleen extended 51 in and liver 11 in below the costal margin Spleen puncture L D bodies present Patient was treated with urea stibamine injected intravenously on successive days from 22 9 32 The following were the doses given (1) 0 lg, (2) 0 lg (3) 0 15 g (4) 0 15 g (5) 0 2 g (6) 0 2 g (7) 0 2 g and (8) 0 2 g Altogether eight injections with a total amount of 13 grms were given. The temperature came down to normal during treatment and no reactions were observed thereafter. At the time of discharge spleen was 11 in below the costal margin and no L D bodies were found on spleen puncture Patient remained in hospital for two months after the completion of treatment

Case No 10—S N M at 30 was admitted on 10 10 32 History of remittent type of fever for 15 days Spleen extended about 2 in below the costal margin and liver was just palpable. Widal was negative. Spleen puncture L D bodies present. Patient was treated with urea stibamine injected intravenously on successive days from 17 10 32. The following were the doses. (1) 0.05 g (2) 0.1 g (3) 0.1 g (4) 0.1 g (5) 0.1 g and (6) 0.1 g Altogether six injections were given with a total amount of 0.55 g. The temperature came down to normal after the third injection and remained so up to the day of discharge from hospital. No reactions were observed during treatment Spleen and liver could not be felt below the costal

margin, and no L D. bodies were found on spleen puncture. Patient remained in hospital for one month after the completion of treatment.

CASE NO 11 -M, æt. 16, was admitted on 19-10-32 History of irregular fever for about 8 months and anæmia with cedema of the lower extremities for about one month. Spleen extended 51 in. below the costal margin. Splech puncture L D. bodies present. Patient was treated with urea stibamine injected intravenously on successive days from 20-10-32 The following were the doses (1) 0 025 g, (2) 0 025 g., (3) 0 05 g, (4) 0 1 g, (5) 0 1 g, (6) 0 1 g., (7) 0 15 g, (8) 0 15 g, (9) 0 2 g and (10) 0 2 g Altogether ten injections were given with a total of 11 grms. The temperature came down to normal after the sixth injection and remained so up to the day of discharge from hospital. No reactions were observed during treatment could just be felt below the costal margin, but no L D bodies were found on spleen puncture and there was neither anæmia nor any œdema of the legs Patient remained in hospital for one month after the completion of treatment

(b) Notes on Cases treated in the Medical Wards of the Campbell Hospital, Calcutta

CASE NO 1—S B, HM, æt 24 History of irregular intermittent fever for nearly 4 months. At the time of admission, liver extended $\frac{3}{4}$ in and spleen $2\frac{1}{2}$ in below the costal arch in the mid-clavicular line. Muscular wasting fairly marked and there was slight pitting on the skin on the tibiæ

Blood Report —R B C —3 900,000, W B C —4,000, polymorphonuclears—61 per cent, lymphocytes—37 per cent, large mononuclears—2 per cent, eosinophiles—nil. Urea stibamine precipitation test strongly positive.

Treatment —Daily injection of urea stibamine 0 05 g I g and 2 g doses eleven injections in all total quantity of urea stibamine given being 1 75 grms

Result — Fever disappeared after the third injection Liver and spleen were no longer palpable. Patient was kept under observation for a period of three weeks after the last injection. At the time of discharge the condition was as follows: Liver and spleen could not be felt below the costal margin. Blood count. R. B. C. —4 000 000. W. B. C. —6 500. polymorphonuclears—68 per cent. lymphocytes—26 per cent. large mononuclears—5 per cent. eosinophiles—1 per cent. Urea stibamine test absolutely negative. Total gain in weight was nearly one stone.

CASE No 2-Dilmaya HF æt 20

Previous History —Irregular intermittent fever for nearly 7 months Patient was cachectic with much muscular wasting anæmia and some cedema of the legs Liver extended 3 in and spleen 5 in below the costal arch in the mid clavicular line

Blood Report —R B C —1 750 000 Hb —30 per cent W B C —2 000 polymorphonuclears—38 per cent lympho cytes—40 per cent large mononuclears—2 per cent eosino philes—nil Formalin and urea stibamine precipitation tests strongly positive

Treatment —Bi weekly injections of urea stibamine from 0.05 to 0.15 g eight injections in all containing one gram of urea stibamine were given preceded by intravenous injections of calcium chloride 10 per cent sol 2 c c each. Eight_injections of hepatrat 3 c c each were also given. As cedema disappeared and the general condition of the patient improved she was given daily injections of urea stibamine thriteen injections in all containing 2.5 grms of urea stibamine.

Result —Fever disappeared after the third injection of urea stibamine Two weeks after the last injection the

condition was as follows Liver was just palpable and spleen, one inch below the costal margin, was hard and appeared to be markedly fibrosed Blood Picture RBC. —3,500,000, WBC—7,000, polymorphonuclears—60 per cent, lymphocytes—36 per cent, large mononuclears—2 per cent, eosinophiles—2 per cent Formalin and urea stibamine tests were negative. Total gain in weight was 1st 10lb.

Case No. 3 —S B Debi, HF, æt 30 Previous history of irregular fever $5\frac{1}{2}$ months, ranging from 102 4° to 97 6° F General condition was poor with much anæmia, dilated heart and general anasarca Liver $2\frac{1}{2}$ in and spleen $4\frac{1}{2}$ in below the costal margin

Blood Report —RBC—1,125,000, Hb—20 per cent, WBC—2,000, polymorphonuclears—62 per cent, lymphocytes—32 per cent, large mononuclears—6 per cent Formalin and urea stibamine tests were strongly positive.

Treatment — The patient was first digitalized and intravenous injections of calcium chloride, 10 per cent sol 2 c.c each, were given for 12 days till the cedema nearly disappeared. Then she was put on daily injections of urea stibamine in 0.05 g., 0.1 g., 0.15 g. and 0.2 g. doses. Altogether fifteen injections were given, amounting to a total of 2.35 grms. of urea stibamine. Fever disappeared after the sixth injection. One week after the last injection, when the patient left hospital, the condition was as follows. Liver was just palpable and spleen ½ in below the costal margin. Blood. Picture: R.B.C.—2,500,000, Hb.—45 per cent, W.B.C.—5,000, polymorphonuclears—65 per cent, lymphocytes—30 per cent, large mononuclears—5 per cent. Formalin and urea stibamine tests were negative. Total gain in weight was 5 lb.

Case No 4.—S N Das, H M, æt 12

Previous History —Irregular intermittent fever for the last 6 months At present the temperature is between 100° and 98° F. Liver $2\frac{1}{4}$ in. and spleen 3 in below the costal

margin at the mid clavicular line Muscular wasting is fairly well marked and so also the black pigmentation of the skin

Blood Report —RBC—2600000 WBC—2400 polymorphonuclears—52 per cent, lymphocytes—38 per cent large mononuclears—8 per cent, eosmophiles—2 per cent Formalin and urea stibamine tests were strongly positive

Treatment —Daily injections of urea stibamine [0 05 g three injections 0 1 g two injections 0 15 g five injections 0 2 g three injections] were given totalling 1 7 grms of urea stibamine

Result —Fifteen days after the last injection blood count showed RBC—3 300 000 WBC—8 000, polymorpho nuclears—56 per cent lymphocytes—32 per cent large mononuclears—2 per cent eosinophiles—10 per cent Urea stibamine test was slightly positive and the spleen was 1½ in below the costal margin. There was no fever. He was given another course of ten injections of 0.2 g of urea stibamine daily. At the end of it the spleen was just palpable at the costal margin and urea stibamine test was negative. The total gain in weight was about half a stone.

(e) Notes on Cases treated in the Out patient Department of Chittaranjan Hospital Calcutta

Case No 1—S HF act 38 came under treatment with history of fever for nearly 6 months and with double rise of temperature for 20 days. Spleen measured 3½ in below the costal arch. Liver was not palpable. Alde hyde test was strongly positive. Patient was given daily intravenous injections of urea stibamine in the following doses (1) 0.05 g (2) 0.05 g (3) 0.1 g (4) 0.1 g (5) 0.1 g (6) 0.15 g (7) 0.15 g and 0.15 g. The number of injections was eight and the total quantity was 0.85 g. The temperature came down to normal after the second injection and remained normal during observation. No reactions were

observed during treatment. Spleen could not be felt below the costal margin and the patient's general condition was satisfactory after the completion of treatment

CASE No 2—M, HM, æt 12, came under treatment with history of fever with double rise of temperature for nearly 5 months. Spleen extended 5½ in below the costal arch and liver was just palpable. Aldehyde test was fairly positive. Patient was given daily intravenous injections of urea stibamine in the following doses (1) 0 05 g, (2) 0 05 g, (3) 0 1 g, (4) 0 1 g., (5) 0 1 g, (6) 0 15 g, (7) 0 15 g, (8) 0 15 g, (9) 0 15 g, and (10) 0 15 g. The number of injections was ten and the total quantity was 1 25 grms. The temperature came down to normal after the third injection and remained so while he was under observation. No reactions were observed during treatment. Spleen was only ½ in below the costal margin and liver was normal. The patient's general condition very much improved one month and a half after the completion of treatment.

CASE No 3—K, H M, æt 17, came under observation with history of continuous fever for about one month and a half Spleen was $2\frac{1}{2}$ in. below the costal arch and liver not palpable Aldehyde test was negative Patient was given daily intravenous injections of urea stibamine in the following doses: (1) 0 05 g, (2) 0 5 g, (3) 0 5 g, and (4) 0 5 g. The number of injections was four and the total quantity was 0.2 g. The temperature came down to normal after the first injection. No reactions were observed during treatment. Spleen could not be felt and patient's general condition was satisfactory.

CASE No 4—S, H F, æt. 12, came under treatment with history of occasional attacks of fever for about one year and with double rise of temperature for 10 days. Spleen measured $3\frac{1}{2}$ in below the costal arch and liver normal. Aldehyde test was fairly positive. Patient was given daily intravenous injections of urea stibamine in the following

doses (1) 0 125 g (2) 0 5 g (3) 0 05 g (4) 0 1 g (5) 0 1 g (6) 0 1 g (7) 0 1 g and (8) 0 1 g The number of injections was eight and the total quantity was 0 625 g The temperature came down to normal after the third injection No reactions were observed during treatment Spleen could not be felt and patient s general condition was very satisfactory

Case No 5—S H M act 5 came under treatment with history of fever for about one year and with double rise of temperature for one month. Spleen measured 4¹ in below the costal arch and liver not palpable. Aldehyde test was strongly positive. Patient was given daily intravenous injections of urea stibamine in the following doses. (1) 0 025 g. (2) 0 025 g. (3) 0 025 g. (4) 0 025 g. (5) 0 025 g. and (6) 0 025 g. The number of injections was six and the total quantity was 0 715 g. The temperature came down to normal after the second injection. No reactions were observed during treatment. Spleen could not be felt below the costal arch and the general condition of the patient improved considerably.

Case No 6—R MM aet 25 came under treat ment with history of fever for about one year and with double rise of temperature for 3 months Spleen measured 6 in below the costal margin and liver was not palpable. Alde hyde test was strongly positive Patient was given daily intravenous injections of urea stibamine in the following doses: (1) 0.5 g (2) 0.1 g (3) 0.1 g (4) 0.1 g (5) 0.1 g (6) 0.1 g (7) 0.10 g (8) 0.2 g (9) 0.2 g and (10) 0.2 g. The number of injections was ten and the total quantity was 1.3 grms. The temperature came down to normal after the third injection. No reactions were observed during treatment. Spleen was about ½ in below the costal arch and the general condition of the patient was very satisfactory.

CASE NO 7—U MM set 25 came under treat ment with history of constant fever for 8 months Spleen

Measured 7 in and liver 2 in. below the costal margin Aldehyde test was fairly positive. Patient was given daily intravenous injections of urea stibamine in the following doses (1) 0 05 g, (2) 0 1 g, (3) 0 1 g, (4) 0 1 g, (5) 0 1 g (6) 0 2 g., (7) 0 2 g, (8) 0 2 g, (9) 0 2 g and (10) 0 2 g. The number of injections was ten and the total quantity was 1 45 grms. The temperature came down to normal after the third injection. No reactions were observed during treatment. Spleen was 2 in. below the costal arch and the liver was not palpable and the general condition of the patient was very satisfactory.

Case No 8—N, MM, æt 10, came under treatment with history of occasional attacks of fever for about one year and with double rise of temperature for two months and a half—Spleen measured 6½ in and liver was not palpable. Aldehyde test was fairly positive—Patient was given daily intravenous injections of urea stibamine in the following doses—(1) 0.5 g., (2) 0.1 g., (3) 0.1 g., (4) 0.1 g., (5) 0.1 g., (6) 0.1 g., (7) 0.1 g., (8) 0.15 g., (9) 0.15 g and (10) 0.15 g. The number of injections was ten and the total quantity was 1.1 grms—The temperature came down to normal after the third injection and remained so during the period of observation—No reactions—Spleen was 2 in below the costal arch and the general condition of the patient was very satisfactory

Case No 9—A B., A I M, æt 28, came under treatment with history of remittent fever for about a month Spleen measured 2 in below the costal arch. Aldehyde test negative and urea stibamine test positive. Patient was given intravenous injections of urea stibamine on successive days in the following doses. (1) 0 05 g, (2) 0'05 g, (3) 0 1 g, (4) 0 1 g, (5) 0'1 g, (6) 0'2 g., (7) 0 2 g. and (8) 0 2 g. The total number of injections was eight, the total quantity being one gram. The temperature came down to normal after the second injection. No reactions.

could not be felt below the costal margin and the general condition of the patient was very much improved

Case No 10—R D H M are 36 came under treatment with history of continuous fever for 13 days. There was no enlargement of the spleen. Aldehyde test negative Widal positive against B typhosus 1/100 L D bodies were found in the thick film from peripheral blood. Patient was given intravenous injections of urea stibamine on successive days in the following doses. (1) 0.05 g. (2) 0.05 g. (3) 0.1 g. (4) 0.1 g. (5) 0.1 g. (6) 0.2 g. (7) 0.2 g. and (8) 0.2 g. The number of injections was eight and the total quantity was one gram. The temperature came down to normal after the fourth injection. No reactions. General condition of the patient was satisfactory.

CASE No 11—A K HF at 18 came under treatment with history of irregular attacks of fever for 9 months. Spleen measured 7 in below the costal arch Aldehyde test strongly positive. Patient was given intra venous injections of urea stibamine on successive days in the following doses. (1) 0.05 g. (2) 0.1 g. (3) 0.1 g. (4) 0.1 g. (5) 0.15 g. (6) 0.2 g. (7) 0.2 g. (8) 0.2 g. (9) 0.2 g. and (10) 0.2 g. The number of injections was ten the total quantity being I.5 gims. The temperature came down to normal after the third injection. No reactions. Spleen could not be felt below the costal margin at the end of the period of observation and general condition of the patient considerably improved.

Case No 12—B H F at 72 came under treatment with history of fever with bronchits for 6 months and hæmoptysis on several occasions. Spleen was 6 in below the costal arch. Aldehyde test strongly positive. Patient wis given intravenous injections of urea stibamine on successive days in the following doses. (1) 0 0.2 g (2) 0 0.2 g (3) 0 0.5 g (4) 0 0.5 g (5) 0 0.75 g (6) 0.075 g (7) 0 1 g (8) 0 15 g (9) 0.2 g and (10) 0.2 g. The number

of injections was ten, the total quantity being one gram. The temperature came down to normal after the eighth injection. No spleen could be felt below the costal margin and the general condition of the patient was satisfactory.

Case No 13—B M, HF, æt 17, came under treatment with history of fever for 15 days with double rise and intermission and with profuse sweating. Spleen measured 2 in below the costal arch. Aldehyde test was positive. Urea stibamine test was strongly positive. Administration of quinine produced no effect. Patient was given intravenous injections of urea stibamine on successive days in the following doses. (1) 0.05 g., (2) 0.05 g., (3) 0.05 g., (4) 0.1 g., (5) 0.1 g., (6) 0.15 g., (7) 0.15 g., (8) 0.15 g., (9) 0.2 g. and (10) 0.2 g. The number of injections was ten and the total quantity was 1.2 grms. The temperature came down to normal after the first injection. No reactions. Spleen could not be felt below the costal margin and the patient's general condition was satisfactory.

CASE No 14-R, A I F, æt 24, came under treatment with a history of periodical attacks of fever and epistaxis for four months Spleen measured 4 in below the costal arch Face was puffy, ankles swollen and patient was anæmic Aldehyde test positive Patient was given daily intravenous injections of urea stibamine in the following doses (1) 0.05 g, (2) 0.05 g, (3) 0.05 g, (4) 0.05 g, (5) 0 1 g, (6) 0 1 g, (7) 0 1 g., (8) 0 15 g, (9) 0 15 g, (10) 0 15 g, (11) 0 15 g, (12) 0 2 g and (13) 0 2 g The number of injections was thirteen and the total quantity was 15 grms The temperature came down to normal during treatment and remained so during the period observation No reactions All the symptoms disappeared. Spleen could not be felt below the costal margin at the end of the period of observation Patient's general condition was satisfactory

CASE No 15 -L MM aet 14 came under treatment with history of fever for about 6 months. There was severe cachexia and the patient could not stand on his legs Spleen extended to the pelvis Aldehyde test strongly positive Patient was given daily infravenous injections of urea stibamine in the following doses (1) 0 025 g (2) 0 025 g (3) 0 025 g (4) 0 025 g (5) 0 05 g (6) 0 05 g (7) 0 05 g, (8) 0 05 g (9) 0 075 g, (10) 0 075 g (11) 0 1 g (12) 0 1 g, (13) 0 1 g (14) 0 1 g (15) 0 15 g and (16) 0 2 g The number of injections was sixteen and the total quantity was 1 2 grms. The temperature came down to normal during treatment and remained so during the period of observation No reactions At the end of the period of observation the spleen could not be felt at the costal margin and his general condition remarkably improved

CASE No 16 -S K MF act 38 came under treat ment with history of fever for 3 months and marked anzemia Spleen palpable on deep inspiration Aldehyde test positive Patient was given daily intravenous injections of urea stiba mine in the following doses (1) 0 05 g (2) 0 1 g (3) 0 15 g (4) 0 15 g (5) 0 15 g (6) 0 15 g (7) 0 15 g (8) 0 2 g (9) 0 2 g and (10) 0 2 The number of injections was ten the total quantity being 1 5 grms. The temperature came down to normal after the first injection. No reactions were observed during treatment Spleen could not be felt on deep inspiration and there was well marked general improve ment of the patient

TABLE SHOWING THE NUMBER OF INJECTIONS AND THE TOTAL AMOUNT OF UREA STIBAMINE IN EACH CASE

	Case No	\	No of injections	Total Amount of Uren Stibnmine
(a)	1 2 3 4 5 6 7 8 9 10	·	10 11 9 12 11 10 10 10 8 6	1 2 g 1 5 g 1 15 g 1 15 g 1 35 g 0 95 g 1 7 g 1 175 g 1 3 g 0 55 g 1 1 g
(b)	1 2 3 4	1	11 13 15 13	175 g 25 g 235 g 172 g
(c)	1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16		8 10 4 8 6 10 10 10 8 8 10 10 10 10 10	0 85 g 1 25 g 0 2 g 0 625 g 0 715 g 1 3 g 1 45 g 1 0 g 1 0 g 1 5 g 1 2 g 1 5 g 1 5 g

OBSERVATIONS

This paper describes a series of cases of kala-azar cured by the intensive treatment with urea stibamine given intravenously on successive days in the following hospitals (1) Carmichael Medical College Hospital, Calcutta, (2) Chittaranjan Hospital, Calcutta, (3) Campbell Hospital, Calcutta and (4) the Out Patient Department Chittaranjan Hospital Calcutta There were no untoward symptoms and temperature generally came down to normal after one or two injections. The number of injections was on an average about ten and the total amount of urea stibamine required in each case is shown in the above table.

We recommend that in suitable cases this intensive treat ment with urea stibamine should be adopted. One remark able feature in this form of treatment was that the total amount of urea stibamine required for a cure in twenty nine out of thirty one cases, was about 1 5 grams or less. This would reduce the cost of treatment to a considerable extent. This fact is of much importance from an economic stand point in the treatment of the disease where its mass treatment has to be taken into consideration by any Government as a prophylactic measure in endemic areas of kala azar.

Reference

Bhm chr Phaldath nd Bhm hii Up danath (1931) Th Inta Anim ni Tr tmentof Kalaza Joun tof Tr pical Med en a d Hygi Aug 15 1931

CAMPAIGN AGAINST KALA-AZAR IN INDIA

- (1) Segregation and removal
- (2) Mass treatment of infected individuals in early days of antimony treatment
- (3) Recent mass treatment with urea stibamine and its success in the campaign
- (4) Dermal Leishmanoid—a possible means of infection
- (5) Conclusions

Compulsory evacuation of the infected areas in Assam (India) was the only available preventive measure against kala-azar before the days of antimonial treatment of the disease, and apparently successful results were obtained by Dodds Price by adopting measures of removal and segregation of infected individuals in certain tea gardens operations were adopted by the Government of Assam, in Golaghat, in 1912 It consisted of removing the patients and their families to a distance not less than 300 yards from the infected house. The infected house was burnt down The valuable properties were disinfected and the less valuable ones destroyed, compensation being paid for their destruction It was, however, observed that removal of the infected family alone was not sufficient to eradicate the infection from a village, as fresh cases of the disease appeared in the neighbouring houses, the year after the removal of the infected family It was, therefore, decided to adopt measures to remove, as "contacts," the families

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who lived in the immediately adjoining houses. The results were more successful but the process had frequently to be repeated year after year as the disease recurred in houses beyond the excised area. In many cases and particularly in areas of old standing infection, it was found that better results followed by moving the whole village at the beginning instead of moving sections of the village in successive years. In this way the mortality was less and the cost remained the same At first these operations seemed to be hopeful and in 1916 it was hoped that if no new factors arose to vitiate the calculations one might look for the extinction of the disease in the areas treated in the above way within a year or two. This hope was not however realised as an epidemic of influenza during the cold weather of 1918 1919 changed the whole situation and there was a recrudescence of the disease in areas where it had been dormant. Further the disease threatened to spread to areas previously uninfected

To control the disease the Government of Assam some time ago adopted certain measures under the provisions of the Epidemic Diseases Act The regulations provided for the notification on the recommendation of the Sanitary Commissioner of any village area found to be infected with kala azar for the prohibition of migration from that area and for the compulsory removal of any of its inhabi tants from an infected site and for the destruction of the infected house and property. The infected families that is to say those in which a case of kala azar was discovered were grouped in an infected camp and contacts. ie their neighbours and any other families which for any reason were under suspicion were located in another group of houses forming the contact camp The remainder were located in a healthy camp which was meant to form the nucleus of a village There was no migration from notified areas and the intercommunication with un

infected villages was greatly limited and chiefly confined to visits between relatives It was found that the removal of a community from an infected area and treatment of those among them who were infected with the disease would terminate the outbreak in that particular community, but the method upon a large scale was prohibitively expensive It was found that inspite of the effective prevention of migration of individuals from infected areas and limitation of communication between infected and healthy villages, a gradual diffusion of the disease took place, and year after year, new infections were discovered in previously uninfected villages to which the same expensive measures of removal and control had to be applied When the general recrudescence which followed the epidemic mentioned above had to be faced, experience showed that segregation on the scale which would be necessary to deal with a widespread prevalence would be prohibitively expensive and administratively impossible (McCombie Young)

Since the treatment of kala-azar by intravenous injection of tartar emetic had been introduced, treatment as a method of prevention was originally put forward by Knowles in 1920 as an alternative to the methods of prevention by segregation The results following mass treatment with tartar emetic were found to be encouraging and early treatment of the first one or two cases seemed to control an outbreak In some cases it appeared to extinguish the disease entirely, perhaps by preventing the establishment of the conditions of the site infection, if the first case had acquired the infection elsewhere In practically all cases the mass treatment seemed to prevent the outbreak from assuming extensive propor-The indications seemed to be that, where only one case came under observation, there was a reasonable chance that no more cases would occur if early treatment -was adopted When several cases were seen for the first time and if they had remained for some time unrecognized and untreated and the opportunities for the establishment of site infection were ample then no amount of trentment of cases seemed to be able to extinguish an outbreak and under those circumstances a perennial crop of cases was to be expected. It seems that early compulsory treatment had a distinct preventive action where it was efficiently applied in the early stages of a village infection but removal to a fresh site was still necessary to terminate an infection when it became deeply rooted by delay in action

The present day treatment campaign against kala azar in Assam has been of immense value as a prophylactic measure. The present day campaign against the disease in Assam is well described in Health Bulletin No. 9 (Government of India Central Publication Branch 1927) containing the Treatment Campaign against Kala azar in Assam as drawn by Major Murison Director of Public Health Assam, and the following extracts are made therefrom—

The treatment of the disease in Assam with tartar emetic began in 1919 when only a comparatively small number of cases were treated. In the special kala azar dispensaries and out centres sodium antimony tartrate manufactured by Messrs Burroughs Wellcome and Company London and put up in soloid form was formerly used exclusively.

Although treatment with this drug was very successful it had the disadvantage of being long and tedious. Treat ment was therefore difficult to enforce as patients who had been completely incapacitated by the disease improved so considerably after a few injections that they discontinued treatment altogether or attended very irregularly. This irregularity made it very difficult to effect complete cures in spite of the regulations in force under the Epidemic Diseases Act to compel patients to undergo a complete course of treatment, the campaign against the disease was

greatly handicapped by the number of patients who were stopping treatment

To overcome this difficulty communiqués were regularly issued inviting the co-operation of the people. Much propaganda work has been done by means of lantern demonstrations and illustrated posters and pamphlets on the disease, emphasizing the grave dangers of stopping treatment before a complete cure has been effected. This had some effect in reducing the "stopped treatment" cases. It was felt that the above difficulties would be still further overcome, if some drug could be introduced, which was not only as efficacious as sodium antimony tartrate, but took a much shorter time to effect a cure

In 1922 the attention of the Government of Assam was drawn to the most brilliant results obtained by Major Shortt, Director, Kala-azar Commission, in the treatment of kala-azar, while working under the auspices of the Indian Research Fund Association at the Pasteur Institute, Shillong, by the use of an aromatic antimonial discovered by me in 1921 and named *Utea Stibamine*, and which I sent to him for trial in cases of kala-azar at the request of Colonel Greig, Director of Medical Research in India (a post now defunct) I had already obtained very satisfactory results in the course of my research in the treatment of the disease (vide my papers on Urea Stibamine in the Treatment of Indian Kala-azar)

In 1927 the campaign against kala-azar continued with unabated vigour and with conspicuous success in Assam, according to a Government Resolution on the Health Report Both the number of cases treated and the number of deaths decreased by slightly over 30 per cent, as compared with the previous year. All districts in which the epidemic was of importance shared in the decrease.

About the middle of this year the universal mass free treatment of the disease in Assam by the use of urea

stibamine (Brahmachari) was introduced throughout the province. This drug effects a cure in a much shorter time than the inorganic salts of antimony which were first success fully employed against the disease and this results in a very much smaller number of cases which give up the treatment before a complete cure is effected. Cases of relapses which are often more obstinate to treatment than original attacks are thus reduced to a minimum.

Major Shortt Director of the Kala azar Commission in India lecturing at the Health Interchange of the League of Nations while describing the campaign against kala azar in Assam considered it to be one of the most successful experiments in public health measures ever adopted and sounded a very strong note of warning against the relaxation of the effort as the disease ceases to have epidemic and again assumes endemic form. He emphasized that it was of the utmost importance to eradicate the endemic foci of infection so that it might not again after a number of years assume an epidemic form. He stated that ' we can only repeat and ask others to repeat that the measures taken by treatment and otherwise within the next few years are likely to be more important and to have a more far reaching effect if resolutely pursued than the most intensive antikala azar measures carried on during the height of the epidemic year

I would now like to point out that in adopting prophy lactic measures one has to guard against possible kala azar carriers as a source of infection. Cases of kala azar with obscure anæmia and cedema, with or without enlargement of spleen and with history of little or no fever may sometimes be found in endemic areas of kala azar and these cases should be properly searched for and properly treated so as to prevent their remaining unsuspected and thereby becoming sources of infection in areas which have apparently been assumed to be completely free from the disease by mass treatment.

Still more difficult are cases of kala-azar previously treated with antimony and apparently cured and which, after one or two years, manifest a remarkable granuloma in the skin containing Leishmania donovani, a condition first discovered by me and named by me as "Dermal Leishmanid". These cases are not so uncommon as I first believed them to be. That being the case, I would suggest that such propaganda should be arranged by the Health Department of infected areas that patients cured of kala-azar might be educated to have their skin carefully examined for the presence of Dermal Leishmanoid for one year or so after completion of treatment. If such lesions are discovered, they should again be treated with a course of urea stibamine

Whatever may be discovered in future about the actual mode of infection, these skin lesions are always infective as they give positive flagellate culture on N.N.N medium. They may propagate the disease by direct contact with the abraded skin of an healthy individual, or if the sandflies are responsible for the propagation of the disease then, as has been recently pointed out by Wenyon, it must be a relatively easy matter for them to take up the parasites from these lesions in the skin as they do in oriental sore.

The following extract from the speech of His Excellency Sir John Kerr, while bidding farewell to the second Legislative Council in Assam (1926), shows the value of the campaign against kala-azar in Assam After referring to the value of the treatment of this disease with urea stibamine, His Excellency said "We may now say that victory, if not in sight, is assured The progress in the campaign against kala-azar in Assam has been phenomenally rapid, and if it continues at the present rate, there is an excellent prospect of the dread scourge being brought under complete control in a few years"

I do not desire to make here any speculative suggestions for the destruction of, or protection against, sandflies as a prophylatic measure since they may be as valueless as any that could have been formulated a few years ago for the destruction of bed bugs which were once supposed to be the carriers of the disease. Instead of asking the poor patient to adopt measures which may be expensive for him to carry out and subsequently turn out to be ineffective one would like to wait till the transmission problem is solved I would now like to listen with very keen interest to any measures that may have been adopted for the eradication of kala azar in the Mediterranean Basin

In conclusion, I would point out that General Gorgas speaking in 1914 on yellow fever control in the Americas stated that its eradication would command the attention and the gratitude of the world and that the thing could be done. To day yellow fever is in full retreat in the Americas. The same will one day be said of kala azar and it may be hoped that before long the disease will be completely banished from India and other parts of the world where it occurs. As I have stated in this paper, the signs of its retreat in Assam are already within sight thanks to the intensive mass treatment of the disease with urea stibamine. When the disease disappears from the world, then one of the highest triumphs of tropical medicine will be achieved.

In my opinion the eradication of kala azar with the recent improvements in its treatment will be a much easier problem than that of malaria

CHEMOTHERAPY OF ANTIMONIAL COMPOUNDS

In studying trypanosomiasis, Ehrlich demonstrated that the trypanosomes assimilated the organic derivatives of arsenic only when the arsenic was present in the trivalent and not in the pentavalent form. Similarly, the experiments of Kolle, Hartoch, Rothermundt and Schurmann have demonstrated that compounds containing pentavalent antimony are not organo-tropic except in large doses and are at the same time slightly parasito-tropic. Preparations containing trivalent antimony were as a rule exceedingly toxic to the organism and at the same time it was also demonstrated by the researches of these observers that for antimony compounds, soluble or insoluble, organic or inorganic, to be of therapeutic value in trypanosomiasis, the antimony must be in the trivalent form

My more recent researches and those of others who have followed me have proved that the pentavalent aromatic antimonials are more potent in the treatment of kala-azar than the trivalent antimonyl tartrates

Metallic antimony is also singular in its chemotherapeutic properties, having the property of being introduced into the veins in the crude form of a fine suspension without any danger of capillary blocking, as has been demonstrated by the observations of Plimmer and Fry as well as those of Ranken in trypanosomiasis and of myself in kala-azar

I shall try to explain what I consider the mechanism by which metallic antimony exerts its parasiticidal properties.

Being an element it does not contain any group or radicals which may complicate any explanation that may be suggested for its action

Levaditi has propounded a general law with reference to all the members of the nitrogen family of elements occupying Group V of Mendeleeff's Periodic Table such as arsenic antimony or bismuth. According to this law these elements exhibit their parasiticidal properties only after they are acted upon by the tissues cg if fresh extract of liver is added to them, they become actively treponemicidal.

Recent unpublished experiments carried on by myself have demonstrated that soon after an intravenous injection of metallic antimony into rats in non toxic doses it disappears from the circulation and appears mostly in the spleen and bone marrow. In the spleen particles of antimony can be found chiefly inside large cells with small nuclei and faintly stained protoplasm. These are probably of the nature of clasmatocyte cells.

It has been observed by Meleney and others that in kala azar these clasmatocyte cells are developed as a tissue reaction out of the reticulo endothehal system. Inside these cells I hold that the antimony is converted into soluble compound containing—Sb=O in a reactive state which exerts its parasiticable action.

The therapeutic action of metallic antimony is greater than that of Sb_2O_3 as well as the antimonyl tartrates because I hold that although these compounds contain the radicle -Sb=O yet it does not exist in them in the reactive stage upon which depends the therapeutic value of an antimonal

The mechanism of the action of metallic antimony when introduced into the vein is therefore that it is first taken up by the cells of the reticulo endothelial system and then converted into a soluble antimony compound containing —Sb=O in the reactive stage (stiboxyl)

By studying the excretion of antimony in man after intravenous injections of a therapeutic antimonial, I have observed that in the case of tartar emetic, the curve of excretion is one slowly converging to the base line valent aromatic antimonials of the type of urea stibamine follow a curve, the first portion of which, representing the excretion during the first 24 hours, is abrupt, similar to what is observed in the case of organic arsenicals, and the second portion follows a curve similar to that found in the case of tartar emetic in which the antimony exists in the trivalent form In other words, after 24 hours or so the organic pentavalent antimonial undergoes a reduction giving rise to a compound, -Sb=O, in the reactive stage This also explains why urea stibamine is more efficacious than either tartar emetic or Sb₂O₃. On the other hand, inorganic antimonates are more or less useless therapeutically as they are excreted as antimonates in the urine without undergoing any change in the body

Generally speaking, the toxicity of the antimonyl tartrates depends upon their antimony content. Notable exceptions are in the case of ammonium antimonyl tartrate and quinine antimonyl tartrate, in which the toxicity is low, especially in the latter. The possibility of using these compounds in therapeutics should therefore be borne in mind. The latter may have the advantage of combining the therapeutic properties of antimony and quinine. Ammonium antimonyl tartrate is the least toxic of all the inorganic tartrates and therapeutically I consider it superior to sodium antimonyl tartrate or tartar emetic.

I have not been able to confirm the observations of Farghar and Gray that the toxicity of the antimony content of quinine antimonyl tartrate is only one-fifth that of tartar emetic, though I agree with them that its toxicity is less than that of tartar emetic. I have confirmed their observations that quinine acid tartrate, on boiling with antimony trioxide,

is converted into the more toxic quino toxin antimonyl tartrate. I have not been able to confirm their conclusions that the sodium salt is less toxic than the potassium salt. I have confirmed Plimmer and Thompson's observations that the lithium salt is more toxic than the sodium or potassium salt and that the toxicity of sodium and potassium salts is equal.

Regarding the excretion of antimony I and my colla borators have observed a law that after single or repeated injections of an antimonial the amount of antimony excreted in the urine during the first twenty four hours is fairly proportional to the amount injected

Further there is a concentrative limit of antimony in the body after repeated injections of tartar emetic and this is the safeguard against any cumulation of this drug in the tissues when the concentration reaches this point

Aromatic Antimonials —Antimonials of the stibeno benzene type have not yet come into use in the treatment of human diseases though they have been used with indifferent results in the case of certain diseases of animals

Phenyl stibinate of Sodium —The minimum lethal dose of phenyl stibinate of sodium is three and a half times less than that of p amino phenyl stibinate of sodium while its maximum tolerated dose is 35 times less. Injected into lower animals it gives rise to a hæmorrhagic nephritis and other symptoms of severe antimony poisoning. This compound has little or no use in therapeutics but the introduction of NH into its benzene nucleus at once diminishes its toxicity and raises its therapeutic value to a remarkable extent.

Acetyl para amino phenyl stibinate of Sodium —By acetylation it is expected on theoretical grounds that the toxicity of p stibanilic acid would be reduced. This acetyl compound has been used in the treatment of kala azar with unsatisfactory results. Besides it becomes toxic with age in

India and it has now come into disuse But I still hold that pure acetyl-p-amino phenyl-stibinate of sodium should again be given a trial in kala-azar and may in future be found to be free from all those toxic effects that were exhibited in the original compound put on the market under the name of stibenyl

Stibamine — As the sodium salt formed after hydrolysis of the acetyl compound corresponds to atoxyl or soamin and is sodium-p-stibanilate, I have named it stibamine Comparing its toxicity with that of the acetyl compound, it will be seen that the introduction of the acetyl group into it does reduce its toxicity as in the case of corresponding arsenic compound. Thus, while in the case of ars-acetin the toxicity is markedly diminished by the introduction of acetyl group into atoxyl (being one-fifth that of atoxyl), in the case of sodium-p-stibanilate and the acetyl compound, my observations have shown that their toxicity is the same, the M.T.D. being 35 grammes per kilo of body weight in guinea-pigs given intramuscularly in the case of both the compounds.

Chloro-acetyl-para-amino-phenyl-stibinate of Sodium — This is a compound formed by the replacement of one hydrogen atom in the benzene nucleus of the acetyl compound by chlorine. I have given it the name of chlorostibacetin. It has been put on the market under the name of stibosan (von Heyden) and it has been claimed that the introduction of chlorine increases its stability, and that it can be stored in ordinary stoppered bottles and weighed out when required and is therefore most useful for general purposes. In my opinion, such a compound has more or less the same stability as the inorganic antimonates and, therefore, there is less chance of the production of the reactive —Sb=O in the tissues after their administration which I consider responsible for the beneficial results following the administration of an antimony compound

Urea stibamine —I consider that urea stibamine owes its therapeutic value to the presence of NH₂CO. The same holds good with the compound which I have named stib glycine amide.

If my theory is correct that the therapeutic value of an inorganic antimonial depends upon the ease with which it can be converted in the tissue into an antimony compound cotaining a radicle -Sb=O in the reactive stage then the same should also apply to the aromatic antimonials

I have given above certain aspects of chemotherapy of antimony in kala azar in respect of which it has been mostly studied If the principle that I have suggested regarding the chemotherapy of antimonial compounds in kala azar infection does not apply to other diseases in which antimony is indicated then the problem must be much more compli cated than what one would at first imagine I would therefore like to hear from my audience here whether the same principle that I have propounded above applies to other diseases as they occur in Egypt such as bilharziasis I have no experience of the treatment of this disease and I would like to listen with very keen interest to the experience of the learned speakers here regarding the chemotherapy of antimony with regard to this disease as well as trypano somiasis

We are just beginning to reach the fringes of the science of chemotherapy in general and this especially holds good in the case of that wonderful element antimony

Some of the compounds referred to here have been mentioned in my paper on *Urea Stibamine* in the treat ment of Indian Kala azar which is to be read at this Congress

UREA STIBAMINE IN THE TREATMENT OF INDIAN KALA-AZAR

My justification in reading this paper is to bring to the notice of the medical profession in Egypt and other parts of the Mediterranean basin the value of an antimony compound which has been of immense benefit in the treatment of Indian kala-azar. I shall begin my remarks by giving here a short history of its introduction in the treatment of kala-azar.

The chemotherapy of aromatic antimonials in kala-azar infection has been the subject of my research for many years In 1920, shortly after I had been financed by the Indian Research Fund Association for carrying on researches into the treatment of kala-azar, acetyl-p-amino-phenyl stibinate of sodium and amino-phenyl-stibinate of sodium were prepared for the first time in India in my laboratory in the Calcutta Campbell Medical School, and I immediately brought to the notice of Government, the Governing Body of the Indian Research Fund Association, and the then Secretary of the Calcutta School of Tropical Medicine, the possibilities of the potentialities of these compounds in the treatment of Indian kala-azar, my conclusions being based on the theoretical grounds, from an analogy of the value of the corresponding compounds of arsenic, namely, ars-acetin and atoxyl in the treatment of certain protozoal diseases

The acetyl compound (=stibacetin, stibenyl) was used more or less successfully outside India in the treatment of kala-azar and other forms of Leishmaniasis (Caronia,

Kharma Marinuchi Spagnoloi) Manson Bahir successfully used it in a case of I ala azar. In India Machie and others found unsatisfactory results in the treatment of kala azar with stibenyl. Early in 1921 I discovered that urea could combine with stibanilic acid and that the resulting compound surpassed all my expectations in its value in the treatment of Indian kala azar. Its introduction and my researches into the chemotherapy of antimonial compounds in kala azar infection under the auspices of the Indian Research Fund Association opened up a new vista in the treatment of the disease in India in face of the untoward results obtained with stibenyl by Mackie and others. This compound I named urea stibanyine.

Until recently tartar emetic or sodium antimonyl tartrate was extensively used for the treatment of I ala azar in India. These have now been completely replaced by urea stiba mine on account of its high therapeutic value and marked superiority over the antimonyl tartrates.

So remarkable was the therapeutic value of this compound that even before the results of my observations were published the medical profession in and outside Calcutta came to recognize its value—soon after its discovery from the reports of cases treated with the drug in my wards in the Campbell Hospital Calcutta—inly first series of successful cases were published in October 1922. In 1923. Shorti and Sen in Assam—reported having obtained more brilliant results with this compound than what was obtained by me in my first series of cases.

The Governing Body of the Indian Research Fund Asso ciation quickly recognized its value from the reports of my cases in Calcutta and also of those obtained from Shortt and other successive Directors of the Pasteur Institute Shillong from Christophers Director of Kala azar Commission who reported from his experience in Assam about its remarkable efficacy from medical officers of tea estates in

Assam, and from the Government of Assam In Calcutta its value was recognized by the physicians of the Calcutta. Medical College Hospitals and successive Superintendents of the Calcutta European Presidency General Hospital In these institutions it was quickly introduced and extensively used with brilliant results. Its reputation quickly spread all over Assam, Bengal, Bihar and Orissa, and to more distant places in India such as Madras, Sanawar, Simla Hills and other places too numerous to mention, and every observer who used the drug was convinced of the great advance made by the discovery of the compound and its introduction in the treatment of Indian kala-azar

Remarking on its therapeutic value, the Secretary, Scientific Advisory Board, Indian Research Fund Association, wrote as follows "Both the Director of the Kala-azar Commission and Lieut -Col E D W Greig consider that this drug has a highly specific action in kala-azar, and its value has been abundantly testified to by those who tried it both in so far as it shortens the period of treatment and in so far as it seems able to cure intractable cases or cases which are resisting the antimony treatment"

I give here some of the important points with regard to the compound

Chemical Constitution and Toxicity—I consider that the chemical constitution of urea stibamine is NH₂ CO C₆H₅ SbO OH ONH₄ containing the group NH₂ CO

Table I

Dose per kg	Number of guinea pigs used	Number died	Remarks
0 7 grm 0 65 ,, 0 6 ,, 0 5 ,, 0 45 ,, 0 4 ,, 0 35 ,	4 2 4 4 4	4 2 2 1 1 1 Nıl	M L D Maj L D M T D

The maximum tolerated dose of urea stibamine per kilo gramme of body weight is twenty three times that of tartar emetic in the case of the guinea pig. The effective dose of urea stibamine in the treatment of kala azar is five sevenths the tolerated dose for the guinea pig. while in the case of tartar emetic it is eight times the tolerated dose for the same animal. Urea stibamine is therefore a much safer antimo nial for use in the treatment of kala azar than tartar emetic or other antimonyl tartrates.

Table II —The value of urea stibamine in the early stage of the disease

	C IM d Coll H sp — B n rdo M C y d auth						P	Hntly T Est t -F te dBn tj		Auth nd M ty						
С	t	2	3	4	5	6	1	2	3	4	5	1	2	1	2	3
Du ton fil n indy Amunt ngm	18	4	120	8	10	21	135	99	120	60	90	7	15	90	150	150
ft whihulting Tilmunt	0 35				03	0 2	1	-			0 95			١	Ľ	0 2
ngm gvn Prodoft t mnt(ndy)	0 35 14		12		0 5 14	0 2	1 .	0 75	0 7 5	0 7 5	0 95 5	0 35 	03	0.5	0 2	0 35 5

It will be seen from the above that urea stibamine cuts short the course of Indian kala azar to a remarkable degree if administered in its early stage

Table III —A series of cases cured with urea stibamine some of which had an intensive course of treatment consisting of multiple injections every 24 hours

D t n of lines (n m nths) Am unt ng m aft which	3	5	5	7	2	3	1	5	!-	- -	-
	0 15	0 85	03	0 65	0 65	0.8	0 4	0 4	0 75	140	65
d P d ft tmntenho	0 35 120	0 35 72	0 3 72	0 75 36	0 85 58	60	0 4 54	0 4 37	1 35 168	1 2 1	15 97
Dy of observat ft t tm nt	36	25					18	33	40	55 2	10

Table IV —A number of cases of Indian kala-azar cured by less than one gramme of urea stibamine

	Cal Med Coll Hosp – Barnardo, VcCay and author					Shill Shortt	ur Inst ong,— , Greig undu	Author and Maity				
Duration of illness (in days)	60	90	10	21	21	240	mths	90				
Amount in grm after which cult neg	0 55	0 65	0 55	09	04	- ;	0 95	0 7	07	0 5	0 6	
Total amount (in grm)	0 55	08	0.7	09	0 55	0 65	0 95	0 7	07	07	0 6	
Period of treatment (in days)	15	14	10	22	14	6	7	5	6	9	5	

In a combined series of 325 published cases, 98 47 per cent of the cases were cured. One of the cases died of extreme asthenia, being admitted at the age of 65 in a moribund condition. In 298 of these cases, proof of cure was microscopic and cultural examination and disappearance of symptoms, and in twenty-seven cases proof of cure was clinical disappearance of the symptoms and subsequent observation of the cases. One case was resistant

Advantages of treatment with Usea Stibamine—(1) The short course occupying only two to three weeks for a complete cure—(2) The appearance of early and marked leucocytosis, rapid disappearance of the fever, of splenic and hepatic enlargement and of anæmia, ædema and cachexia, in fact, of all the symptoms of the disease, and reversion to normal state of health. (3) The absence of symptoms of intolerance after its administration—(4) It is most valuable in the treatment of relapses or in the cases resistant to sodium antimonyl tartrate or tartar emetic. (5) Observations have shown that early cases may be cured after four or five injections

Indications and Contra-indications in the treatment of Kala-azai with Urea Stibamine —(1) Urea stibamine is indicated in all cases of kala-azai. It has been used in cases

complicated with bronchitis or dysentery without any un toward results. In cases of Leishmania dysentery, it cures the condition. If, however dysentery develops during treat ment then it is desirable to stop the injection for a time or give it in smaller doses. (2) In eases with marked nephritis and ædema, begin with smaller doses. If the ædema in ereases give the doses at longer intervals than before. (3) Very advanced cases should be treated by beginning with small do es which should be slowly increased.

Urea stibamine has been observed to manifest no deterioration or other changes either in physical and chemical characters or in therapeutic properties if kept in sealed ampoules under ordinary conditions

Among the other therapeutic antimonials that have been discovered by me for use in the treatment of Indian kala azar in the course of my research are (I) stibamine (2) chloro stibacetin (3) stibgly eine amide (=N phenyl glyeine amide p stibinate of sodium) (4) the glucose derivatives of these compounds and (5) stib heetine

I shall not refer to any of these compounds besides point ing out that some of them have been found to be of thera peutic value in the treatment of Lala azar especially stibgly cine amide

In 1921, while discussing with the Director of the Calcutta School of Tropical Medicine about the therapeutic value of urea stibamine soon after its discovery I drew his attention to the possibility of obtaining therapeutic aromatic antimonials from the Chemische Fibrik von Heyden Stibosan and neo stibosan are among von Heyden's preparations that have since been used in the treatment of Indian kala azar

Of all aromatic antimonials that have been used up to the present time in India the most extensive trial has been given to urea stibinine. The published reports consist of cases treated by different observers and under different conditions.

and are therefore most valuable; it has stood the test of time for upwards of seven years.

A review of the treatment of Indian kala-azar with urea stibamine is given in my Treatise on Kala-azar (John Bale, Sons and Danielsson Ltd., London, 1928), to which I would refer my audience for a detailed account of the compound

I would now quote the remarks of Shortt, Director of Kala-azar Commission, and Sen (1925) regarding the therapeutic value of urea stibamine "We consider that the value of urea stibamine has been established as the most efficient drug at present in use in the treatment of Indian kala-azar". This statement remains equally true to-day. To this may be added the remarks of Dodds Price "I am of opinion that urea stibamine is a most valuable remedy in the resistant types of the disease, and I strongly urge that it should be resorted to if, after a few injections of sodium antimonyl tartrate, a patient does not show marked improvement."

Gentlemen, antimony is a wonderful element. Centuries ago, Valentine, who wrote about it in The Triumphant Chariot of Antimony, did so with awed devotion. He said "He who writes of antimony needs a great consideration and a most ample mind. One man's life is too short to be perfectly acquainted with all its mysteries." Centuries later, it was supposed to do so much harm that the graduates in Medicine in the University of Heidelberg had to swear never to use antimony. And to-day it has been found to be a specific in one of the most terrible of tropical diseases.

THE CONQUEST OF KALA AZAR

[This is the first part of the author s Presidential Address Section of Medical Research | I wenty fifth Session of the Indian Science Congress Calcutta 1938 The second part on Certain Obser Lotions on the Chemotherapy of Wolaria is published in the Second Volume]

LADIES AND GENTLEMEN,

We are meeting under the most tragic circumstances. The sudden death of Lord Rutherford who was to be our General President has cast the deepest gloom over our Congress. The world has lost in him one of its most distinguished scientists who smashed the atoms and though he breathes no more his discoveries may one day enable one to travel to that mysterious region among the ntoms and molecules wherein enters the breath of life. We express our deepest sense of sorrow to Lady Rutherford and the bereaved family. We are fortunate in having another most famous scientist as our General President and we have every reason to believe that under the guidance of Sir James Jeans, this momentous meeting of the Indian Science Congress will come to a successful issue.

The earliest epidemic of knln near in Bengal occurred in the seventies of the last century, when it was probably complicated with malarin. As I stated elsewhere in this epidemic it was noted by a contemporary writer that countries that once smiled with peace health and prosperity had been turned into hot beds of disease misery and death and that the fell disease had mocked every human effort and absorbed

in its powerful grasp, day by day and inch by inch, every blessed spot which once used to be prized for its salubrity... This was the old Buidwan fever

In more recent times the epidemic of the disease in Nowgong district of Assam produced such an appalling mortality that there was a decrease of 31.5 per cent in the population of the place in the decade 1891-1900

The mortality from the disease has now been reduced from 90% or more to 1 or 2 per cent — Including complicated cases, it has been reduced from 99 to less than 10 per cent

The conquest of kala-azar may be said to have begun when Cristina and Caronia obtained remarkable results in infantile kala-azar of the Mediterranean basin by the use of tartar emetic. Rogers introduced this drug into India for the treatment of Indian kala-azar and obtained similarly satisfactory results. Soon after the introduction of tartar emetic the speaker introduced sodium antimonyl tartrate for its treatment. This was taken up by others, as the compound was considered to be more powerful and less toxic than tartar emetic.

The next step in the treatment of the disease was the introduction by the speaker of intravenous injection of metallic antimony in a state of fine subdivision as an impalpable powder

It has been observed (Brahmacharı and co-workers) that when metallic antimony is injected intravenously in a state of fine subdivision, the particles are picked up by the same cells in the spleen as those that harbour the parasites of kala-azar and that in the struggle that ensues the fight ends most remarkably in the complete destruction of the parasites in the speediest way (See Plate facing page 226)

A special outfit has been devised for the intravenous injection of metallic antimony

The advantage of intravenous injection of metallic antimony is that the number of injections generally required is not more than three or four to bring about a complete cure It is one of the most powerful leishmanocides and may be tried in cases in which other antimonials have failed. The great objection to its use is the complicated technique of the operation of injection which is a serious obstacle in the mass treatment of the disease.

Although treatment with tartar emetic or sodium antimonyl tartrate was very successful in the treatment of kala azar it was found that in the campaign against the disea e it had the disadvantage of being long and tedious. In Assam which was once the hot hed of the disease treatment was therefore found difficult to enforce as patients disconti nued treatment altogether or attended very irregularly after a few injections This irregularity made it very difficult to effect a complete cure The Director of Public Health Assam once noted that in spite of the regulations in force under the Epidemic Diseases Act to compel patients to undergo a complete course of treatment the campaign against the disease was greatly handicapped by the large number of patients who stopped treatment. To overcome this difficulty communiques were regularly issued inviting the co operation of the people. Much propaganda work was done by means of lantern demonstrations and illustrated posters and pamphlets on the disease emphasising the great dangers of stopping treatment before a complete cure was effected Though this had some effect in reducing the Stopped Treatment cases, still such cases continued to exist and it was felt that the difficulties in reducing the number of such cases would be overcome more effectively if some drug could be introduced which would be more efficacious than tartar emetic and take a much shorter time to effect a cure

The introduction of certain organic compounds of antimony in the treatment of kala azar infection has been the subject of the speaker's research for many years and in 1920 some of them were prepared for the first time in India in his laboratory in the Calcutta Campbell Hospital, and he wrote to the Indian Research Fund Association that the potentialities of the preparation of these compounds in India would in future be as great as those of cinchona plantation

Early in 1921, the speaker discovered an urea antimony compound for the treatment of kala-azar. Its introduction and his other researches on antimonial compounds opened up a new vista in the treatment of the disease by means of therapeutic organic antimonials, just as the discovery of salvarsan led to the introduction of organic arsenicals in the treatment of spirochætal diseases. This urea compound was named 'urea stibamine'. Soon after its discovery the author suggested to the Director of the Calcutta School of Tropical Medicine the possibility of obtaining therapeutic antimonials from Von Heyden and this led to their introduction into this institution for the treatment of kala-azar.

The first series of cases treated with urea stibamine were published early in 1922. Soon after this, remarkable results were obtained with it by Shortt in Shillong to whom the compound was sent for trial at the instance of the Director of Medical Research appointed by the Government of India. The value of this compound was quickly recognized and it was introduced, after an experimental trial, by the Government of Assam for the mass treatment of kala-azar.

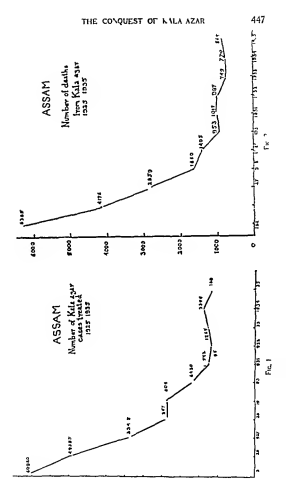
I now proceed to demonstrate the value of urea stibamine in the campaign against kala-azar in Assam as obtained from statistics from the Annual Public Health Reports of the Government of Assam for the years 1925-35 and the Government Resolutions thereon. This drug has been more or less successively in use by the Government of Assam for over twelve years and to-day it is in universal use in the province. For some time experiments with neostibosan were conducted side by side with urea stibamine. The use of neostibosan was subsequently discontinued.

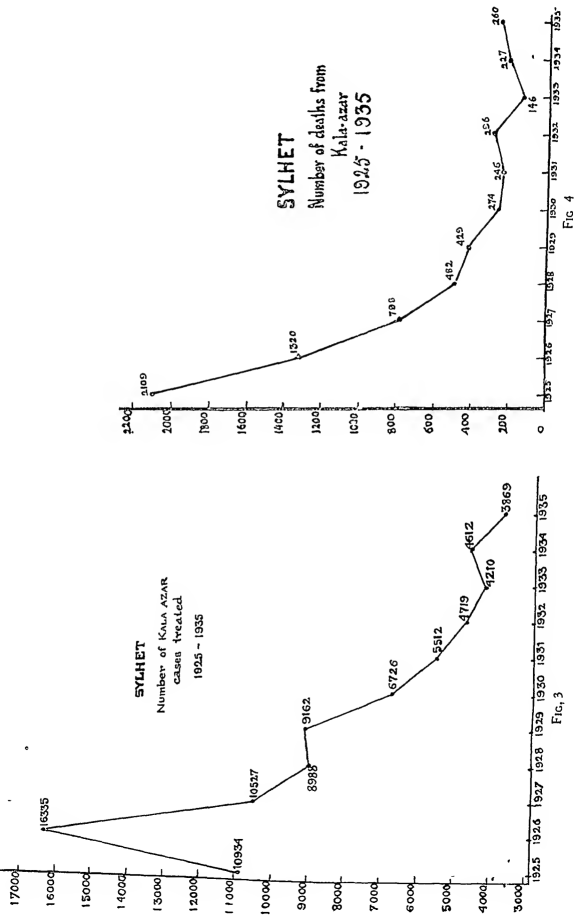
In their resolution on the Annual Public Health Reports

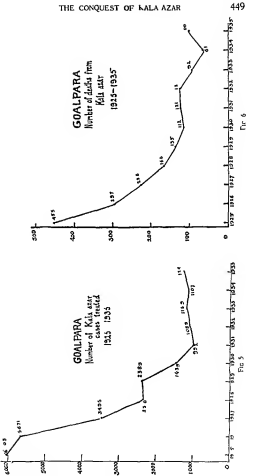
of the Province of Assam for the year 1926 the Govern ment of Assam noted that the treatment with urea stibamine proved very successful and there was a very satisfactory decrease of over 1 000 in the number of 'Stopped Treat ment cases

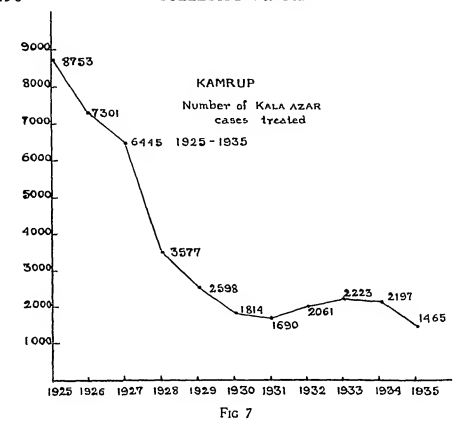
Figure 1 gives the number of cases of kala azar treated in Assam as a whole from 1925 to 1935 showing a marked fall in the incidence of the disease from 60 940 in 1925 to 11 100 in 1935 Figure 2 gives the death rate from kala azar in Assam as a whole from the year 1925 to 1935 showing a marked fall from 6 365 in 1925 to 845 in 1935 gives the number of cases of kala azar treated in the district of Sylhet during the same years showing a marked fall in the incidence of the disease from 10 934 in 1925 to 3 869 in 1935 Figure 4 gives the death rate from the disease in the district of Sylhet from the year 1925 to 1935 showing a marked fall from 2 109 in 1925 to 260 in 1935 Figure 5 gives the number of cases of kala azar treated in the district of Goalpara during the same years showing a marked fall in the incidence of the disease from 6 003 in 1925 to 1 245 in Figure 6 gives the death rate from the disease during the same years in Goalpara showing a marked fall from 453 in 1925 to 100 in 1935 Figure 7 gives the number of cases of the disease treated in the district of Kamrup during the same years showing a marked fall in the incidence of the disease from 8 753 in 1925 to 1 465 in 1935 Figure 8 gives the death rate for the same years in the district of Kamrup from the year 1925 to 1935 showing a marked fall from 1 120 in 1925 to 176 in 1935 Figure 9 gives the number of cases treated in the district of Darrang during the same years showing a marked fall in the incidence of the disease from 5 262 in 1925 to 738 in 1935 Figure 10 gives the death rate for the same years in the district of Darrang for the years 1925 to 1935 showing a marked fall from 478 in 1925 to 91 in 1935 Figure 11 gives the number of cases

showing a marked fall in the incidence of the disease from 13,895 in 1925 to 1,651 in 1935. Figure 12 gives the death rate from the disease for the same years in the district of Nowgong showing a marked fall from 1,445 in 1925 to 52 in 1935. Figure 13 gives the number of cases treated in the Garo Hills during the same years showing a marked fall in the incidence of the disease from 1,952 in 1925 to 690 in 1935. Figure 14 gives the death rate from the disease for the same years in the Garo Hills showing a marked fall from 435 in 1925 to 58 in 1935.









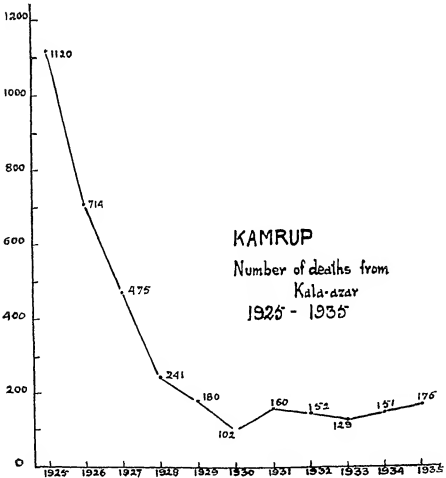
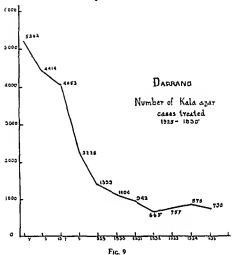
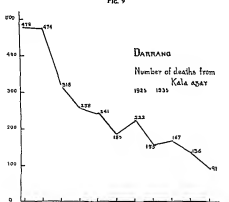
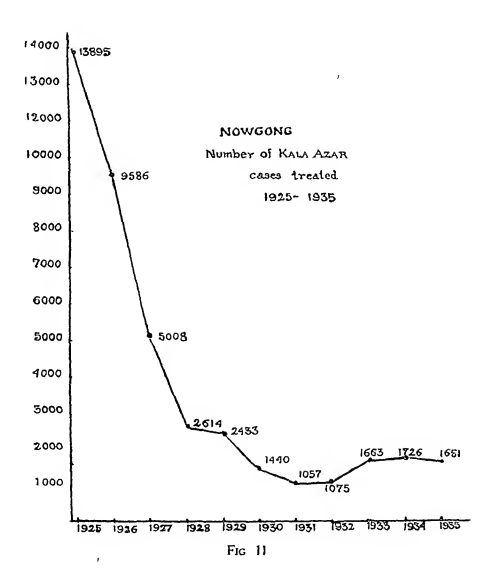
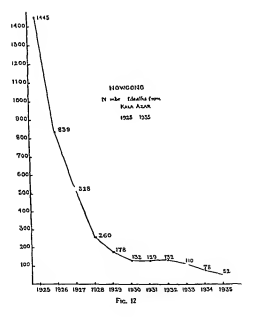


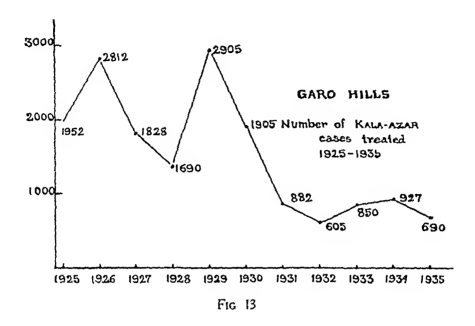
Fig 8

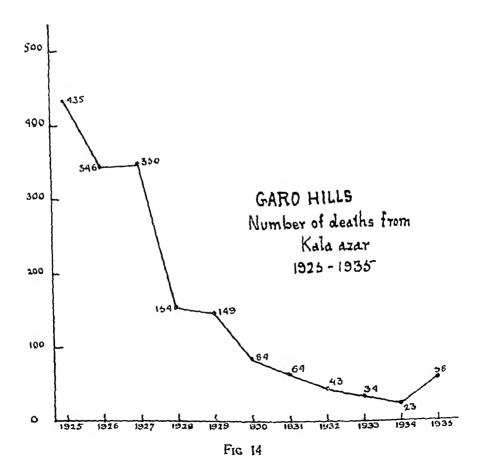












<u>3</u>

88

4 612 227

5%

STATISTICS OF THE NUMBER OF KALA AZAR CASES TREATED BY THE GOVERNIENT OF ASSAM AND THE NUMBER OF DEATHS FROM THE DISEASE

	161	22.2	
	1930	6726	
Sulh t	10.29	62† 291 6	
	1928	8 948	
	1261	10 527	
	1976	16 335	
	1725	10 934 2 109	

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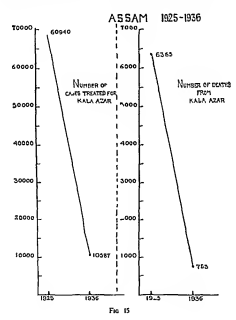
No of cas No of d 1 >

			1,,,	
	1932	1 245	8	
	1934		52	
-	1933		26	
	1932		1 089	
	1631		262	
	0691		1439	
Goalpa a	u261	-	2 389	-
	9761	- -	2316	_
	161		3 495	İ
	9,61		5 671	:
	1925		6 003	3
	, ,		N f c fr ted	0

Kamrup

Year	1925	1926	1927	1928	1929	1930	1661	1932	1933	1934	1935
No of cases treated No of deaths	8,753	7,301	6,445	3,577	2,598	1,814	1,690	2,061	2,223	2,197	1,465
1					Darrang				_		_
Year	1925	1926	1927	1928	1929	1930	1931	1932	1933	1934	1935
No of cases treated No of deaths	5,262	4 4 1 4 4 7 4	4,053	2,228	1,399	1,106	942	665 155	757	876	738
					Nowgong	_			_		
Vest	1025										
1001	(76)	1926	1927	1928	1929	1930	1631	1932	1933	1934	1035
No of cases treated No of deaths	13,895 1,445	9,586 839	5,008	2,614	2,433	1,440	1,057	1,075	1,663	1,726	1.651
					Garo H.ll.			701	011	78	52
Vooy	201				1						
1 5 4 1	6761	1926	1927	1928	1929	1930	1831	1932	1933	1934	1025
No of cases treated	1,952	2.812	1 828	1,000	- -		- -				567
No of deaths	435	346	350	154	2,905	, 1,905 84	882 64	605	850	927	069
							_	<u> </u>	ξ.	23	58

The number of cases treated and of deaths in 1936 could not be incorporated in the above figures and tables, as the Annual Health Report for 1936 was not available when the figures and tables were printed off Fig 15 gives a comparative statement of the number of cases treated and deaths in the Province of Assam from 1925 to 1936



The figures for Cachar which are not exhibited in the above diagrams are very interesting. Out of 5 188 cases 58-767B

showing a percentage of less than 1.08 per cent. Out of 574 cases treated in 1936 the number of deaths was 2 showing a percentage of less than 0.3 per cent.

It will be seen from the above figures and tables that the effect of the treatment with urea stibamine during the above-mentioned years on the incidence of kala-azar in Assam and its mortality has been phenomenal. The disease has lost all its terrors in the province and people who suffer from kala-azar are less afraid than those who suffer from malaria.

The Kala-azar Commission, India, used, throughout the seven years of its existence, urea stibamine only, in the routine treatment of kala-azar. According to them the acute fulminating type characteristic of the peak period of an epidemic responded to treatment with urea stibamine extraordinarily promptly and with an almost dramatic cessation of fever, diminution in size of the spleen and return to the normal condition of health. It may be expected that similar beneficial results will be obtained in other epidemics of the disease.

In 1933, the Director of Public Health, Assam, noted that 'urea stibamine was our mainstay in the treatment of kalazar' 'Since 1923, when reliable figures for the diseases first became available to the end of the year under report, no less than 328,591 persons have been brought under treatment. It is no exaggeration to say that approximately 3 25 lacs of valuable lives have been saved to the Province.'

Sir John Kerr, once Governor of Assam, in his farewell address to the Assam Legislative Council referring to the value of urea stibamine stated that 'the progress in the campaign against kala-azar in Assam has been phenomenally rapid and if it continues at the present rate there is an excellent prospect of the dread scourge being brought under complete control in a few years' This has now come to

pass as you have just seen from the statistics quoted in the present address and as was once predicted by the Director of Public Health Assam one day we shall be successful in stamping the disease out of the province. The same may also be said of other parts of the world where the disease occurs. The discovery of a powerful specific for the disease its limited distribution and rarity of relapses lend support to this conclusion.

A few antimonials have been tried intramuscularly in the treatment of kala azar. Among these may be mentioned sodium N phenyl glycine amide 4 stibinate (antimony ana logue of tryparsamide) and sodium sulphomethylene stibani late (antimonyl analogue of sulph arsenol) which have been successfully used by the author while neostibosan has been used with success by Napier

In studying the treatment of kala azar one finds that the enemy in the process of destruction tries to retreat from the internal organs to some other parts of body just as it is noticed in the case of human warfare the conquered foe tries to hide himself in hills and jungles to elude the pursuit of the con queror. This was first discovered in 1922 when the speaker observed certain remarkable skin eruptions caused by Leishmania donovani developing in kala azar patients two or three years after completion of antimonial treatment and apparent cure though under ordinary conditions in kala azar the skin shows ve y little involvement or none. On ginally considered very rare these skin lesions have been subsequently found not to be an uncommon condition.

The disease was named dermal leishmanoid by the speaker when first discovered and subsequently called dermal leishmaniasis in the Carmichael Hospital for Tropical Diseases attached to the Calcutta School of Tropical Medicine

The various types of the disease will not be described here in detail

For the photograph of the first recorded case of dermal leishmanoid See Plate LXIII, facing page 52

The photograph of a case of annular variety of the disease and of the photomicrographs of sections of skin showing the presence of leishmania-laden cells just under the epidermal layer and of leishmania-laden pigment-carrying cells in the superficial layer of the dermis are reproduced here

Viable leishmania have been cultured from these skin lesions in test tubes and sandflies. They are therefore a source of infection and the conquest of kala-azar cannot be regarded complete unless these lesions are either averted or quickly cured. Not infrequently, they require a prolonged course of antimonial treatment and some of them are very resistant and may be dangerous carriers of antimony-resistant parasites. The author has recommended combined treatment with intravenous injection of urea stibamine and inunction of metallic antimony.

It is evident that in the campaign against kala-azar and its conquest, proper handling of cases of dermal leishmanoid is an important point to be taken into consideration

The constitution of urea stibamine has been a matter of some controversy. As pointed out by Gray and coworkers it is the most interesting of therapeutic antimonials. Originally considered by the speaker to be a urea salt of para-amino-phenyl stibinic acid, it was subsequently described by him to be ammonium carbamino-stibanilate. More recently Gray and co-workers have studied the chemical constitution and physiological properties of the compound carefully and exhaustively in an important paper in the Proceedings of the Royal Society (1931). They have shown that urea stibamine is disubstituted urea consisting mostly of S-diphenylcarbamide-4 · 4'-distibinic acid as its active organo-metallic constitution, containing some amount of protective colloids to make it water-soluble. Its constitution is therefore, according to these observers,

different from that of compounds of the type of neostibosan or neostam which are salts of para aminophenyl stibinic acid

The conquest of kala azar by campaign against the disease by treatment of the affected individuals is from what I have shown one of the most remarkable feats in chemotherapy. Whether a prophylactic dose of urea stibamine to persons living in kala azar infected areas just like the prophylactic use of quinine in malaria will be of any value or whether an inunction of metallic antimony may be recommended to be used by them as a routine practice is a matter for investigation.